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(71) Applicant (for all designated States except US): CHI-RON CORPORATION [US/US]; 4560 Horton Street, Emeryville, CA 94608-2916 (US).

(72) Inventors; and

60/465.273

60/465.535

60/468,312

60/473,144

60/495,024

60/505,652

60/510,781

60/529,464

60/536,177

60/560.757

(75) Inventors, and
Riccomparison (for US only): RAPPUOLI,
Rine [IT/IT]; c/o Chiron Corporation, P.O. Box 8097,
Emeryville, CA 94662-8097 (US). MASIGNANI,
Vega [IT/IT]; c/o Chiron Corporation, P.O. Box 8097,
Emeryville, CA 94662-8097 (US). STADLER, Konrad [DE/DE]; c/o Chiron Corporation, P.O. Box 8097,
Emeryville, CA 94662-8097 (US). GREGERSEN,
Jens-Peter [DE/DE]; c/o Chiron Corporation, P.O.

Box 8097, Emeryville, CA 94662-8097 (US), CHIEN, David [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). HAN, Jang [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). POLO, John [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). WEINER, Amy [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). HOUGHTON, Michael [GB/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). SONG, Hyun, Chul [KR/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). SEO, Mi, Young [KR/US]; c/o Chiron Corporation, P.O. Box 3097, Emeryville, CA 94662-8097 (US). DONNELLY. John, J. [US/US]; c/o Chiron Corporation, P.O. Box 3097, Emeryville, CA 94662-8097 (US). KLENK, Hans, Dleter [DE/DE]; c/o Chiron Corporation, P.O. Box 3097. Emeryville, CA 94662-8097 (US). VALIANTE, Nicholas [US/US]; c/o Chiron Corporation, P.O. Box 3097, Emeryville, CA 94662-8097 (US).

(74) Agents: HALE, Rebecca, M. et al.; Chiron Corporation, Intellectual Property R338, P.O. Box 8097, Emeryville, CA 94662-8097 (US).

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(54) Title: THE SEVERE ACUTE RESPIRATORY SYNDROME CORONAVIRUS

(57) Abstract: An outbreak of a virulent respiratory virus, now known as Severe Acute Respiratory Syndrome (SARS), was identified in Hong Kong, China and a growing number of countries around the world in 2003. The invention relates to nucleic acids and proteins from the SARS coronavirus. These nucleic acids and proteins ran be used in the preparation and manufacture of vaccine formulations, diagnostic reagents, kits, etc. The invention also provides methods for treating SARS by administering small molecule antiviral compounds, as well as methods of identifying potent small molecules for the treatment of SARS.

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THE SEVERE ACUTE RESPIRATORY SYNDROME CORONAVIRUS

All documents cited herein are incorporated by reference in their entirety.

RELATED APPLICATIONS, FROM WHICH PRIORITY IS CLAIMED

This application incorporates by reference in its entirety US provisional patent application 60/462,218, Attorney Reference No. PP20474.001, filed on April 10, 2003 via Express Mail with the US post office, US provisional patent application 60/462,465, Attorney Reference No. PP20480.001, filed on April 11, 2003 via Express Mail with the US post office, US provisional patent application 60/462.418, Attorney Reference No. PP20480.002, filed on April 12, 2003 via Express Mail with the US post office, US provisional patent application 60/462,748, Attorney Reference No. PP20480.003, filed on April 13, 2003 via Express Mail with the US post office. US provisional patent application 60/463,109, Attorney Reference No. PP20480.004, filed on April 14, 2003 via Express Mail with the US post office, US provisional patent application 60/463.460, Attorney Reference No. PP20480.005, filed on April 15, 2003 via Express Mail with the US post office, US provisional patent application 60/463,668, Attorney Reference No. PP20480.006, filed on April 16, 2003 via Express Mail with the US post office, US provisional patent application 60/463,983, Attorney Reference No. PP20480.007, filed on April 17, 2003 via Express Mail with the US post office, US provisional patent application 60/463.971. Attorney Reference No. PP20480.008, filed on April 18, 2003 via Express Mail with the US post office, US provisional patent application 60/464,899, Attorney Reference No. PP20480.009, filed on April 22, 2003 via Express Mail with the US post office, US provisional patent application 60/464,838, Attorney Reference No. PP20507.001, filed on April 22, 2003 via Express Mail with the US post office, US provisional patent application 60/465,273, Attorney Reference No. PP20518.001, filed on April 23, 2003 via Express Mail with the US post office, US provisional patent application 60/465,535, Attorney Reference No. PP20518.002, filed on April 24, 2003 via Express Mail with the US post office, US provisional patent application 60/468,312, Attorney Reference No. PP20480.010, filed on May 5, 2003 via Express Mail with the US post office, and US provisional patent application 60/473,144, Attorney Reference No. PP20480.011, filed on May 22, 2003, US provisional patent application 60/495,024, Attorney Reference No. PP20480.012, filed on August 14, 2003 via Express Mail with the US post office, US provisional patent application 60/505,652, Attorney Reference No. PP20480.013, filed on September 23, 2003 via Express Mail with the US post office, US provisional patent application 60/510,781, Attorney Reference No. PP20480.014, filed on October 11, 2003 via Express Mail with the US

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FIELD OF THE INVENTION

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The invention relates to nucleic acids and proteins from Severe Acute Respiratory

Syndrome (SARS) Virus. These nucleic acids and proteins can be used in the preparation and
manufacture of vaccine formulations for the treatment or prevention of SARS. The invention
also relates to diagnostic reagents, kits (comprising such reagents) and methods which can be
used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The
invention also relates to methods for the treatment or prevention of SARS utilizing small
molecule viral inhibitors and combinations of small molecule viral inhibitors and kits for the
treament of SARS.

BACKGROUND OF THE INVENTION

An outbreak of a virulent respiratory virus, now known as Severe Acute Respiratory Syndrome (SARS), was identified in Hong Kong, China and a number of other countries around the world in 2003. Patients typically had symptoms including fever, dry cough, dyspnea, headache, and hypoxemia. Isolates of the SARS virus appear to have homology with at least the RNA polymerase gene of several known coronaviruses. A phylogenetic analysis of this homology is presented in Peiris et al., "Coronavirus as a possible cause of severe acute respiratory syndrome", Lancet, published online April 8, 2003 at http://image.thelancet.com/extras/03art3477web.pdf, incorporated herein by reference in its entirety. Other sequenced fragments of the SARS virus genome appear to overlap with the open reading frame 1b of coronaviruses. See, Drosten et al., "Identification of a Novel Coronavirus in Patients with Severe Acute Respiratory Syndrome", New England Journal of Medicine, published online at http://www.nejm.org on April 10, 2003, incorporated herein by reference in its entirety.

The Genome Science Center in British Colombia, Canada published on its website (http://www.bcgsc.ca/bioinfo/SARS/) a draft genome assembly of 29,736 base pairs of a virus believed to be a SARS virus, referred to as the TOR2 isolate. This draft genome assembly is given herein as SEQ ID NO: 1.

The Centers for Disease Control (CDC) published a nucleotide sequence of a SARS-CoV strain (SEQ ID NO: 2) on its website (http://www.cdc.gov/ncidod/sars/pdf/nucleoseq.pdf). The CDC

has also published a phylogenetic tree of the predicted N, S and M proteins (attached as FIGURE 6). This tree places the SARS virus outside any of the previously known coronavirus groups.

There is a growing need for prophylactic or therapeutic vaccines against the SARS virus as well as diagnostic and screening methods and compositions to identify the presence of the virus in, e.g., mammalian tissue or serum.

SUMMARY OF THE INVENTION

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The invention relates to nucleic acids and proteins from Severe Acute Respiratory-Syndrome (SARS) virus. These nucleic acids and proteins can be used in the preparation and manufacture of vaccine formulations for the treatment or prevention of SARS. Such vaccine formulations may include an inactivated (or killed) SARS virus, an attenuated SARS virus, a split SARS virus preparation and a recombinant or purified subunit formulation of one or more SARS virul antigens. Expression and delivery of the polynucleotides of the invention may be facilitated via viral vectors and/or viral particles.

The invention also relates to diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention further includes non-coding SARS viral polynucleotide sequences, SARS viral sequences encoding for non-immunogenic proteins, conserved and variant SARS viral polynucleotide sequences for use in such diagnostic compositions and methods.

The invention further relates to vaccine formulations comprising one or more SARS virus antigens and one or more other respiratory virus antigens. Additional respiratory virus antigens suitable for use in the invention include antigens from influenza virus, human thinovirus (HRV), parainfluenza virus (PIV), respiratory syncytial virus (RSV), adenovirus, metapneumovirus, and rhinovirus. The additional respiratory virus antigen could also be from a coronavirus other than the SARS coronavirus. Preferably, the additional respiratory virus antigen is an influenza viral antigen.

The compositions of the invention may further comprise one or more adjuvants. Adjuvants suitable for use in the invention include mucosal, transdermal or parenteral adjuvants. Mucosal adjuvants suitable for use in the invention include detoxified bacterial ADP-ribosylating toxins, such as E. coli heat labile toxoids (e.g., LTK63), chitosan and derivatives thereof, and non-toxic double mutant forms of Bordetella pertussis toxoids. Parenteral adjuvants suitable for use in the invention include MF59 and aluminum or aluminum salts.

The invention also provides methods for treating SARS by administering small molecule compounds, as well as methods of identifying potent small molecules for the treatment of SARS.

In one aspect of the invention a method of identifying a therapeutically active agent is provided comprising: (a) contacting the therapeutically active agent with a cell infected with the SARS virus; (b) measuring attenuation of a SARS related enzyme.

In a more particular embodiment, the therapeutically active agent is a small molecule. In another more particular embodiment, the therapeutically active agent is a nucleoside analog. In another more particular embodiment the therapeutically active agent is a peptoid, oligopeptide, or polypeptide. In another embodiment the SARS related enzyme is SARS protease. In another embodiment the SARS related enzyme is SARS polymerase. In still another embodiment the SARS related enzyme is a kinase. Methods of identifying therapeutically active agents for treatment of SARS virus infection are further discussed in Section V below.

In another aspect of the invention a method of treating a human infected with SARS is provided comprising administering a small molecule to a patient in need thereof. In one embodiment the small molecule is an inhibitor of SARS protease. In another embodiment the small molecule is an inhibitor of SARS polymerase. In another embodiment the SARS related enzyme is a kinase. In still another embodiment the small molecule is administered orally or parenterally.

The invention also provides the use of such small molecules in the manufacture of a medicament for the treatment of severe acute respiratory syndrome.

Small molecule compounds of the present invention include those of less than 1000 g/mol, preferably with an aromatic region and greater than one heteroatom selected from O, S, or N. Preferred small molecules include, but are not limited to acyclovir, gancyclovir, vidarabidine, foscamet, cidofovir, amantidine, ribavirin, trifluorothymidine, zidovudine, didanosine, zalcitabine, and combinations thereof. Interferons may also be used for treating patients, including interferon-α and interferon-β. Interferon treatment has shown promise in treating SARS in monkeys (Enserink (2004) Science 303:1273-1275), particularly when pegylated (Haagmans et al. (2004) Nature Medicine 10:290-293).

One aspect of the present invention relates to methods for identifying individuals exposed to, and biological samples containing SARS virus (SARSV), and to kits for carrying out the methods. Such methods can utilize nucleic acid detection techniques such as PCR, RT-PCR (the Coronaviridae are RNA viruses), transcription-mediated amplification (TMA), ligase chain reaction (LCR), branched DNA signal amplification assays, isothermal nucleic acid sequence based amplification (NASBA), other self-sustained sequence replication assays, boomerang DNA amplification, strand-displacement activation, cycling probe technology, or combinations of such amplification methods. Such nucleic acid detection techniques utilize oligonucleotides having nucleotide sequence similar to, or complementary to, the SARS viral genome, as primers (e.g., for amplification) and as probes (e.g., for capture or detection), as is well known in the art.

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Alternatively, or in addition to the nucleic acid detection methods described supra, the methods of the present invention can utilize various immunoassay techniques for detection of SARSV antigens and/or antibodies.

Accordingly, the present invention relates to methods of identifying individuals exposed to SARSV, or biological samples containing SARSV, by detecting the presence of SARSV antigens using antibodies which specifically bind to the same. The antibodies are preferably monoclonal antibodies. Quantification of the amount of viral antigens present in a sample of an individual may be used in determining the prognosis of an infected individual. Preferably, the SARSV antigens to be detected are generally one of the structural proteins, particularly those present on the surface of the viral particles and include, for example, the spike glycoprotein (S), also called E2; the envelope (small membrane) protein (E), also called SM; the membrane glycoprotein (M), also called E1; the hemagglutinin-esterase glycoprotein (HE); also called E3; and the nucleocapsid phosphoprotein (N). In preferred embodiments, the antigens to be detected are the S, E and M proteins using antibodies to the same.

The present invention relates to kits for identifying individual SARSV and reagents used in such kits. The kits comprise a first container which contains antibodies which specifically bind to a SARSV antigen and a second container which contains the SARSV antigen. The antibodies are preferably monoclonal antibodies. The kits may be adapted for quantifying the amount of antigen in a sample of an individual. Such information may be used in determining the prognosis of an infected individual.

The present invention relates to methods of identifying individuals exposed to SARS virus, or biological samples containing SARSV, by detecting the presence of antibodies against SARS virus antigen in a sample using SARS antigen. Quantification of the amount of anti-SARS protein from SARS antibodies present in a sample of an individual may be used in determining the prognosis of an infected individual. Any one or more of the viral proteins (structural proteins or nonstructural proteins) may be used as antigen to detect the SARSV antibodies; preferably a SARSV antigen that is conserved amoung SARSV isolates is preferred. In this regard, nonstructural protein (e.g., Pol, Hel, 3CLp, MP, PLP1, PLP2) may be particularly useful.

The present invention relates to kits for identifying individuals exposed to SARS and reagents used therein. The kits comprise a first container which contains antibodies which were produced in response to exposure to an antigen from SARS virus and a second container which contains the SARS antigen(s). The kits may be adapted for quantifying the amount of anti-SARS antibodies present in a sample of an individual. Such information may be used in determining the prognosis of an infected individual.

The present invention relates to methods of identifying individuals exposed to SARS virus, or biological samples containing SARSV, by detecting the presence of nucleic acid from SARS

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virus. Quantification of the amount of SARS nucleic acid present in a sample of an individual may be used in determining the prognosis of an infected individual. The methods utilize oligonucleotide probes and/or primers that are similar or complementary in sequence to the SARSV genome or transcription or replication products. Preferred probes and primers are described herein. Also included in the present invention are kits for carrying out the methods of detecting the SARSV nucleic acid.

The invention further includes a method for the treatment and/or prevention of SARS through the administration of a therapeutically effective amount of at least one antiviral compound from among those described in the US Patents and published international patent applications listed in Table 1 and Table 2. In one embodiment of the method, the antiviral compound is a small molecule. In another embodiment, the antiviral compound is a protease inhibitor. In a further embodiment, the antiviral protease inhibitor is a 3C-like protease inhibitor and/or a papain-like protease inhibitor. In another embodiment, the antiviral compound is an inhibitor of an RNA-dependent RNA polymerase. In another embodiment, a first antiviral compound which is a protease inhibitor is administered with a second antiviral compound which is an RNA-dependent RNA polymerase inhibitor. The invention further provides for the administration of a steroidal anti-inflammatory drug in combination with at least one antiviral compound, for example, from the antiviral compounds described in the documents listed in Table 1 and Table 2.

The invention further provides for a method for the treatment and/or prevention of SARS through the administration of a therapeutically effective amount of at least one antiviral compound from among those described in the US Patents and published international patent applications listed in Table 1 and Table 2 by inhalation. In one embodiment of the method, the antiviral compound is a small molecule. In another embodiment, the antiviral compound is a protease inhibitor. In a further embodiment, the antiviral protease inhibitor is a 3C-like protease inhibitor and/or a papain-like protease inhibitor. In another embodiment, the antiviral compound is an inhibitor of an RNA dependent RNA polymerase. In another embodiment, a first antiviral compound which is a protease inhibitor is administrated with a second antiviral compound which is an RNA-dependent RNA polymerase inhibitor. The invention further provides for the administration of a steroidal anti-inflammatory drug in combination with at least one antiviral compound, for example, from the antiviral compounds described in the documents listed in Table 1 and Table 2 by inhalation. The steroidal anti-inflammatory drug may be administered by inhalation for a local effect or administered for systemic absorption such as via an oral or intravenous route.

The invention further provides the use of an antiviral compound, as defined above, in the manufacture of a medicament for the treatment of severe acute respiratory syndrome.

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The invention further provides for a kit for use by a consumer for the treatment and/or prevention of SARS. Such a kit comprises: (a) a pharmaceutical composition comprising a therapeutically effective amount of at least one antiviral compound from among those described in the US Patents and published international patent applications listed in Table 1 and Table 2 and a pharmaceutically acceptable carrier, vehicle or diluent; (b) a container for holding the pharmaceutical composition; and, optionally; (c) instructions describing a method of using the pharmaceutical compositions for the treatment and or the prevention of SARS. The kit may optionally contain a plurality of antiviral compounds for the treatment of SARS wherein the antiviral compounds are selected from 3C-like protease inhibitors and papain-like protease inhibitors. In a further embodiment, the kit contains an antiviral compound which is an RNA-dependent RNA polymerase inhibitor. When the kit comprises more than one antiviral compound, the antiviral compounds contained in the kit may be optionally combined in the same pharmaceutical composition.

An additional aspect of the invention provides for the use of at least one of the antiviral compounds described in the US Patents and published international patent applications listed in Table 1 and Table 2 for the manufacture of a medicament for the treatment or prevention of SARS.

BRIEF DESCRIPTION OF THE DRAWINGS

- FIGURE 1: Schematic of coronavirus genome organization.
- 20 FIGURE 2: Schematic of coronavirus ORF1a/ORF1b gene products.
 - FIGURE 3 (A C): Alignment of coronavirus polynucleotide sequences for selected genes (including nucleocapsid (N), matrix (M), and hemagluttinin-esterase (HE)).
 - FIGURE 4 (A F): Alignment of coronavirus polypeptide sequences (including ORF1a/ORF1b, nucleocapsid (NP), hemagluttinin-esterase (HE), envelope (Sm or E), matrix (M), and spike (S).
- 25 FIGURE 5: Alignment of spike (S) polypeptide sequences, taken from Figure 4, in the region of the junction of the S1 and the S2 domains, and protease cleavage site for selected coronaviruses.
 - FIGURE 6: CDC phylogenetic tree of SARS-CoV strain (Clustalx 1.82, neighbor-joining tree). Figure 6A shows coronavirus N protein analysis, Figure 6B shows coronavirus S protein analysis, and Figure 6C shows coronavirus M protein analysis.
- 30 FIGURE 7: Conserved and specific sequence of the SARS virus. Figures 7A-7D show multiple sequence alignments (CLUSTAL W 1.82) of the structural proteins of the SARS virus genome (7A: PEP4 Spike protein; 7B: PEP7 small membrane protein; 7C: PEP8 matrix glycoprotein; 7D: PEP13 nucleocapsid protein), which have counterparts in all or some of the other known coronaviruses. Figures 7E-7H show dendrograms reporting the protein distances among the

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sequences in alignments 7A-7D. <u>Labels</u> 229E: human coronavirus; MEV: murine hepatitis virus; TGV: transmissible gastroenteritis virus; AIBV: avian infectious bronchitis virus; BOVINE: Bovine coronavirus; PEDV: porcine epidemic diarrhea virus.

- FIGURE 8: Alignment of the 5'UTR of several coronaviruses, to show consensus nucleotide sequence at the 5'UTR.
- FIGURE 9: Sequences of preferred primers for amplification of the 5'UTR. F and R denote forward and reverse PCR primers, and the numbers indicate nucleotide positions withing Figure 8.
- FIGURE 10: Alignment of the 3'UTR of several coronaviruses, to show consensus nucleotide sequence at the 3'UTR.
- FIGURE 11: Sequences of preferred primers for amplification of the 3'UTR. F and R denote forward and reverse PCR primers, and numbers indicate nucleotide positions within Figure 10.
- FIGURE 12: Coiled-coil prediction for SEQ ID NO: 6042, using Coils program (Figure 12A) or LearrCoil (Figure 12B).
- 15 FIGURE 13: Example of insertion of a reporter gene-of-interest at a site between exisiting SARS , virus genes. Small nonstructural gene products are not depicted schematically.
 - FIGURE 14: Schematic depicting representative examples of SARS virus replicons. Small nonstructural gene products are not depicted schematically.
- FIGURE 15: SARS virus nsp2 proteinase (3CLp) and identification of catalytic and substrate
 20 sites.
 - FIGURE 16: alignment of SARS virus nsp2 proteinase (3CLp) with that of avian IBV, MHV, and BCoV. Residues in dotted boxes are key residues the substrate sites (F, Y & H); residues in solid boxes are catalytic cysteine (C) and histidine (H) residues.
- FIGURE 17: Genome organization of SARS coronavirus. Replicase and structural regions are
 shown, along with the predicted products of cleavage within ORF1a and ORF1b. The position of
 the 5' RNA leader sequence (L), the 3' poly(A) tract and the ribosomal frame-shift consensus
 between ORF1a and ORF1b are also indicated. Each box represent a protein product. They are
 shaded according to the level of amino acid identity with corresponding proteins of other
 coronaviruses (see also Table 2). The SARS-specific genes are white. Positions of the 9 SARSspecific six-base IG sequences (5'-ACGAAC-3'; SEQ ID NO 7293) are indicated by arrows.
 - FIGURE 18: Genome organization of Coronaviruses representative of group 1 (HCoV-229E, accession number: AF304460), group 2 (mouse hepatitis virus MHV, accession number: NC_001846), group 3 (avian infectious bronchitis virus AIBV, accession number: NC_001451)

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and SARS coronavirus. Other completely sequenced coronaviruses used in this study are available at the following accession numbers: porcine epidemic diarrhea virus (PEDV), AF353511; transmissible gastroenteritis virus (TGV), NC_002306; Bovine coronavirus (BCoV): AF220295. Red boxes represent group-specific genes. The position of the leader RNA sequence and poly(A) tract is also indicated in genomes where they are reported. The position of specific IG sequences is indicated by circles of different shades. In the SARS genome, we also find three IG sequences specific for group 2 coronavirus.

FIGURE 19: Topological model predicted for the spike protein anchored to the viral membrane. Structural and predicted functional domains are indicated. The N-terminal region (S1) is predicted to contain the receptor binding domain. Two coiled coil regions within the S2 domain, partially superimposed to leucine zipper motifs are presumably involved in oligomerization. The hydrophobic domain is responsible for membrane anchoring.

FIGURE 20: Phylogenetic tree obtained from the multiple sequence alignment of a 922 bp internal region of the *pol* gene from 12 coronaviruses and SARS. Numbers at the nodes represent the result of a bootstrap analysis and strongly support the branches. Sequences not available within the complete coronavirus genomes have been retrieved from GenBank at the following accession numbers: hemagglutinating encephalomyelitis virus of swine (PHEV), AF124988, Human OC43 virus (OC43), AF124989, canine coronavirus (CCV), AF124986, feline infectious peritonitis virus (FIPV), AF124987, turkey coronavirus (TCV), AF124991, syaloacryoadenitis virus of rats (SDAV). AF124990.

FIGURE 21: 21A. Unrooted tree obtained from the alignment of consensus sequences of the

group I and group II S1 domain of spike proteins (G1_cons and G2_cons) with those of a group 3 spike (AIBV) and the spike of SARS virus. The number indicates the result of a bootstrap analysis. The sequences used to generate the consensus profile from group 1 are: HcoV-229E, accession number P15423; porcine epidemic diarrhea virus (PEDV), acc no: NP_598310; transmissible gastroenteritis virus (TGV), acc no: NP_058424; Canine coronavirus (CCV), acc no: S41453; porcine respiratory virus (PRV), acc no: S24284; feline infectious peritonitis virus (FIPV), acc no: VGIH79. The sequences used to generate the consensus profile from group 2 are: mouse hepatitis virus (MHV), acc no: NP_045300; Bovine coronavirus (BCoV), acc no: NP_150077; Human coronavirus OC43, acc no: P36334; hemagglutinating encephalomyelitis virus of swine (PHEV), acc no: AAL80031; for group 3, only the sequence of the spike protein of avian infectious bronchitis virus (AIBV), acc no: AAO34396 was used. 21B: Schematic representation of cysteine positions in S1 domains of group 1, 2 and 3, compared to the SARS spike. Horizontal bars represent the S1 amino acid sequences (in the case of SARS and AIBV) or the consensus profiles (generated from group 1, G1. cons, and from group 2, G2. cons). The

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length of the bars are not to scale. Relative cysteine positions are indicated by rectangle bars. Only cysteines perfectly conserved within each consensus are reported. Lines connect cysteines conserved between the SARS S1 domain and the consensus sequences as shown.

FIGURE 22: illustration of a Neisseria Adhesin A protein (NadA).

5 FIGURE 23: Raw translation from SARS coronavirus genome (reading frame +1).

FIGURE 24: Raw translation from SARS coronavirus genome (reading frame +3)

FIGURE 25: 1b and Spike open reading frames, separated by *.

FIGURE 26: SARS growth in vero cells.

FIGURE 27: chromatogram of the capture step of SARS coronavirus on Matrix Cellufine

Sulfate Superformance 150/10. Analysis was on 100ml coronoavirus harvest. The left Y axis shows absorbance at 280nm. The right Y axis shows the gradient (%B). The X axis shows the volume (ml).

FIGURE 28: Silver-stained MCS chromatography fractions. Lanes are: (1) marker;

- (2) coronavirus vero cell harvest; (3) coronavirus vero cell harvest, after 0.65μm filtration;
- 15 (4) flowthrough; (5) wash; (6) 20% peak (virus peak). Lanes were loaded with 1 μg of test protein.
 - FIGURE 29: Western Blot of MCS chromatography fractions. Lanes are as described for Fig.28.
 - FIGURE 30: Linear density gradient ultracentrifugation, 15-60% sucrose (SW28, 2 hours, 20000 rpm). The graph shows protein concentration (*),
- FIGURE 31: Silver-stained density gradient fractions on NuPage 4-12% Bis-Tris-Ge (Novex), reduced conditions, heated for 10 minutes at 70°C. Lanes are: (1) marker, (2) 20% peak MCS;
 - (3) density gradient fraction 11; (4) density gradient fraction 12; (5) density gradient fraction 13;
 - (6) density gradient fraction 14; (7) density gradient fraction 15; (8) density gradient fraction 16;
- (9) density gradient fraction 17. The bulk of proteins was in fractions 15 to 17. Lanes 2, 8 and 925 were loaded with 1µg protein.
 - FIGURE 32: Chromatogram of the Capture Step of SARS coronavirus on MCS. Details are as for Figure 27, except that 200ml harvest was used.
 - FIGURE 33: Silverstain (left) and Western Blot (right) of chromatographic fractions. Lanes are as described for Figures 28 and 29, except that lane (6) is the 5% peak. Treatment before
- SDS-PAGE was at room temperature for 30 minutes.
 - FIGURE 34: Density Gradient Ultracentrifugation, 15-40% sucrose (SW28, 2 hours, 20000 rpm). The graph shows protein concentration (*).

FIGURE 35: Silverstain (left) and Western Blot (right) of Density Gradient Ultracentrifugation fractions on NuPage 4-12% Bis-Tris-Ge (Novex), reduced conditions. Lanes are: (1) marker; (2) density gradient fraction 6; (3) density gradient fraction 7; (4) density gradient fraction 8; (5) density gradient fraction 9; (6) density gradient fraction 10; (7) density gradient fraction 15.

- 5 Fractions 7-10 (lanes 3-6) contained pure coronavirus proteins. The bulk of impurities was in fraction 15 (lane 7). Lanes 2, 8 and 9 were loaded with ~1μg protein. Treatment before SDS-PAGE was at room temperature for 30 minutes.
 - FIGURE 36: EM pictures of Density Gradient Fractions 8-10. Figure 36A shows fraction 8; Figure 36B shows fraction 9; Figure 36C shows fraction 10.
- 10 FIGURE 37: Spike/NadA fusion constructs.
 - FIGURES 38 and 39: Results of the expression in *E.coli* of S1_L, S1_L-NadA and S1_L-NadA_{Annehor}. Figure 38 shows SDS-PAGE analysis of total lysates from BL21(DE3)/pET, BL21(DE3)/pET-S1_L and BL21(DE3)/pET-S1_L-NadA_{Annehor}. The bands are indicated by an arrow, and the three lanes are, from left to right: BL21(DE3)/pET; BL21(DE3)/pET-S1_L; BL21(DE3)/pET-
- S1_L-NadA_{Aaachor}. Figure 39 shows (39A) SDS-PAGE and (39B) western blot analyses of total lysates from BL21(DE3)/pET, BL21(DE3)/pET-S1_L-NadA (grown under un-induced condition) and BL21(DE3)/pET-S1_L-NadA (grown under induced condition). The bands are indicated by an arrow, and lanes are, from left to right: BL21(DE3)/pET; BL21(DE3)/pET-S1_L-NadA; BL21(DE3)/pET-S1_L-NadA. The western blot shows the presence of oligomeric forms of the protein.
 - FIGURE 40: Schematic of SARS Spike clones.

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- FIGURE 41: Transient Expression of SARS Spike Proteins (western blot of COS7 cell lysate). Each lane of the 4-20% TG SDS gel was loaded with $20\mu g$ cell lysate (total 1.2mg). The labeling antibodies are shown.
- 5 FIGURE 42: Western blot analyses of COS7 cell lysates on 4% TG SDS gel showing oligomerization state of intracellular S molecules.
 - FIGURE 43: Western blot analyses of COS7 cell lysates on 4-20% TG SDS gel showing Transient Expression of SARS Spike Proteins. Lanes are: (1) mock, AF; (2) mock, DF; (3) nSh, AF; (4) nSh, DF; (5) nSh Δ TC, AF; (6) nSh Δ TC, DF. Each lane was loaded with 5μ l of each sample, 400 μ l total. The blot was labeled with antibody against the His-tagged protein.
 - FIGURE 44: Western blot analyses of COS7 cell medium on 4-20% TG SDS gel showing Transient Expression of SARS Spike Proteins. Truncated spike protein is secreted. Spike proteins were purified from the culture medium (from a 10cm plate), first by a ConA column and then finally by His-tag Magnetic beads. Each lane was loaded with one third of the material.

FIGURE 45: Western blot analyses of COS7 cell lysates on 4-20% TG SDS gel showing glycoslation of SARS spike proteins. In the two left-hand blots (lanes 1-5), samples were boiled in SDS and β-mercaptoethanol; in the two right-hand blots (lanes 6-11), samples were in SDS only, with no boiling. Lanes 1-8 were labeled with a monoclonal raised against the His-tag protein; labes 9-11 were labeled with rabbit anti-SARS antibody.

FIGURE 46: Effect of SARS spike protein expression on cell viability.

FIGURE 47: Western blot analyses of COS7 cell lysates on 4% TG SDS gels showing – oligomerization state of intracellular spike molecules. Blots were labeled with anti-His-tag mAb. The membrane fraction of COS7 cell lysate was fractionated by a sizing column before loading the lanes. Fractions 7 to 14 show bands with kDa values of: 71000, 1400, 898, 572, 365,232, 148 and 99, respectively.

FIGURE 48: Fractionation of cells into aqueous and detergent fractions.

FIGURE 49: Schematic of constructs for use in OMV preparation.

FIGURE 50: SARS HR1 and HR2 constructs.

15 FIGURE 51: Vaccine protection froms SARS in Balb/c mouse model.

FIGURE 52: Expressed on Spike protein in transfected 293 cell lysates (52A) or COS7 cell culture supernatants (52B). Proteins were separated on 4-20% TG SDS gels. The label was anti-His-tag, except for the right-hand three lanes of 52B, where the label was rabbit anti-SARS serum. In Figure 52A, the left-hand three lanes were treated with DTT and were boiled, but neither treatment was used for the right-hand three lanes. In Figure 52B, no DTT was used, but all lanes were heated to 80°C for 5 minutes

FIGURE 53: Western blot of Spike proteins expressed in COS7 cells. Proteins were incubated at room temperature (RT), 80°C or 100°C to check for any effect on molecular weight. FIGURE 54 shows similar experiments on SARS virions.

25 FIGURE 55: Results of a pulse chase experiment, showing expression and processing of SARS spike protein following infection with alphavirus replicon particles. Cells were treated with or without EndoH as shown.

FIGURE 56: Effect of heating on Spike protein trimers.

FIGURE 57: Coomassie blue-stained gel of yeast-expressed proteins. Lanes are: 1-See Blue Standard (10μl); 2-pAB24 gbl (20μg); 3-SARS Spike S1 c.1 gbl (20μg); 4-SARS Spike S1 c.2 gbl (20μg); 5-See Blue Standard (10μl); 6-pAB24 ip (5μl); 7-SARS Spike S1 c.1 (5μl); 8-SARS Spike S1 c.2 (5μl).

FIGURES 58 to 64: Schematics of preparation of yeast expression constructs.

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FIGURES 65 to 66: Yeast-expressed sequences for Spike.

FIGURE 67: Western blots showing expression of SARS spike protein from alphavirus replicon particles and replicon RNA. Figure 67A was run under non-reducing conditions and at room temperature (i.e. no heating), with lanes: (1) VEE/SIN-spike infection; (2) VEE/SIN-GFP infection; (3) Replicon-spike RNA transfection; (4) Replicon-GFP RNA transfection. Figure 67B

was run with SARS virions at different temperatures, as shown.

FIGURE 68: induction of antibody responses in mice. Vaccine groups are: (1) Inactivated SARS

Virus; (2) Truncated Recombinant Spike Protein; (3) Full length Spike: DNA+DNA.PLG+ Alphavirus; (4) Full length Spike: Alphavirus particles only.

10 FIGURE 69: Binding of human monoclonal antibody S3.2 to purified truncated Spike protein. The X-axis shows antibody concentration, and the Y-axis shows ELISA absorbance. The interpolation result is 2158.13.

FIGURE 70: Geometric mean ELISA titers of antibodies induced by the SARS-CoV spike protein delivered as different vaccines (left to right: inactivated virus; $3\mu g$ truncated spike protein; $75\mu g$ DNA encoding truncated spike protein.

FIGURE 71: Neutralization titers after immunization with (left) nSd Δ TC protein or (right) DNA encoding nSd Δ TC, delivered on PLG.

FIGURE 72: Correlation between the spike antigen binding and neutralizing antibodies

FIGURE 73: Western blot of CHO cell lines expressing Spike protein in full-length form (left) or in truncated form (right). Proteins were separated by 4-12% SDS-PAGE, with boiling in DTT and staining by polyclonal serum.

FIGURE 74: Structural components of SARS-CoV spike glycoprotein and expression construct. L denotes leader peptide (residues 1-13), TM the transmembrane, and Cy the cytoplasmic tail segments. The hexa-His tags are not shown.

25 FIGURE 75: Western blot analysis of SARS spike proteins expressed in COS7 cells. In Figure 75A, COS7 cells were transfected with indicated plasmid constructs and the expressed proteins in cell lysates 48 hr post-transfection were analysed by SDS-PAGE (4-20% polyacrylamide) in reducing and denaturing conditions, with proteins visualized by anti-histidine Mab. In Figure 75B, proteins were collected from cell culture medium 48 hr post-transfection and purified first by a ConA column and then by His-tag magnetic beads. Purifed proteins were analysed by SDS-

by a ConA column and then by His-tag magnetic beads. Purified proteins were analysed by SDS-PAGE (4-20% polyacrylamide) and were visualized by anti-SARS rabbit serum.

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FIGURE 76: Endo H sensitivity of C-terminal truncated spike protein (S Δ) found in cell lysate (lanes 1,2) and culture medium (lanes 3,4). Positions of internal S Δ protein and secreted S Δ protein are marked with arrow heads.

FIGURE 77: Oligomeric status of the SARS spike protein. Recombinant S protein oligomer in COS7 cells transfected with the full-length spike construct (nSh). The cell lysates were treated with DTT and/or heat as indicated above each lane. The different forms of S protein in treated and untreated samples were visualized by SDS-PAGE (4% polyacrylamide) and Western blot analysis using anti-histidine MAb.

FIGURE 78: Effect of heat denaturation on the oligomeric status of recombinant S protein in the absence of DTT. The COS7 cell lysates were heated before the electrophoresis as indicated and the S proteins were visualized as described fogiFigure 77.

FIGURE 79: Effect of heat denaturation on the oligometic status of spike protein in SARS virion particles. SARS-CoV were grown in Vero cells, purified and solubilized from the virion particles by SDS, heat-denatured as indicated and visualized as described in Figure 77, except that rabbit antiserum against the purified virus was used as a probe.

FIGURE 80: Analysis of the oligomeric status of SARS virion spike protein by cross-linking experiment. Solubilized SARS virion proteins were treated with DMS. Both untreated (–) and DMS treated (+) virion proteins were heat denatured in the absence of DTT and visualized by 4% PAGE followed by silver staining.

20 FIGURES 81 & 82: Analysis of the oligomeric status of truncated spike protein by heat denaturation. Truncated spike protein within COS7 cell lysates (81) or secreted into culture medium (82) were heat denatured as indicated in the absence of DTT and visualized by Western blot analysis.

FIGURE 83: Reactivity of deglycosylated full-length spike oligomer with conformational and non-conformational antibody. The full-length recombinant spike oligomer was partially deglycosylated with PNGase F in non denaturating condition and visualized by Western blot analysis using anti-histidine Mab (lane 1,2,3) or rabbit antiserum against purified SARS CoV (lane 4,5,6).

FIGURE 84: Localization of expressed SARS spike proteins in fractionated COS7 cell lysate visualized by western blot. Cells were transfected with indicated plasmids and lysed with Dounce homogeniser in hypotonic buffer 48 hr post transfection. Cell lysate was centrifuged to obtain soluble cytosol and insoluble membrane fraction that was further solublized by 4% Triton X-100. Proteins were heated with SDS at 80 C and analysed by SDS-PAGE (4-20% polyacrylamide) in reducing condtion. Proteins were visualized by anti-histidine Mab. The

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cytosol fractions were loaded in lanes 1, 3, and 5 and the membrane fractions were loaded in lanes 2, 4, and 6.

FIGURE 85: Intracellular and surface expression of recombinant full-length (A,D) or truncated (B,E) spike protein in COS7 cells. The cells were fixed at 48 hrs posttransfection and either treated with detergent (Cytofix/perm, BD Biosciences) for intracellular immunofluorescence (A,B,C) or with 2% paraformaldehyde for cell surface immunofluorescence observation (D,E,F) at x40 magnification. Mock transfected cells (C,F) were included as controls.

FIGURES 86-105: SDS-PAGE od *E.coli* expressed proteins. Tot = total protein; Sol = soluble protein fraction. Labels are protein names (Tables 26-30).

10 FIGURE 106: Immunofluorescence after administration of vector encoding optimsed N antigen.

FIGURE 107: Immunofluorescence of (A) native and (B) codon-optimsed M sequences.

FIGURE 108: Immunofluorescence of (A) native and (B) codon optimsed E sequences.

FIGURES 109-111: Western blots of Vero cells using rabbit antibodies obtained after immunization with spike proteins expressed in *E.coli*.

15 FIGURE 112: Spike protein expression in 293 cells. Lanes: (M) Markers; (1) Mock transfected; (2,6) cells expressing nS protein, lysate; (3,7) cells expressing nSdTC protein, lysate; (4,8) cells expressing nS protein, supernatant; (5,9) (4) cells expressing nSdTC protein, supernatant. Staining antibody: (2 to 5) mouse serum obtained after DNA immunization; (6 to 9) rabbit serum obtained after immunization with whole killed virus.

20 FIGURE 113: Six reading frames of SEQ ID NO: 9968.

FIGURE 114: Six reading frames of SEQ ID NO: 10033.

FIGURE 115: Alignment of bovine coronavirus pol 1ab (top row; SEQ ID NO: 10068), avian infectious bronchitis pol 1ab (second row; SEQ ID NO: 10069), murine hepatitis virus pol 1ab (third row; SEQ ID NO: 10070), SEQ ID NOs: 9997/9998 (fourth row) and a consensus sequence (bottom row; SEQ ID NO: 10071).

FIGURE 116: Schematic of coronavirus genome organization.

FIGURE 117: Schematic of coronavirus ORF1a/ORF1b gene products, including "*" region.

FIGURE 118: Alignment.

FIGURE 119: Alternative start codons within SEQ ID NO: 10080.

30 FIGURE 120: Six reading frames of SEQ ID NO: 10084.

FIGURE 121: Alignment of SEQ ID NO: 10033 and SEO ID NO: 10084.

FIGURE 122: Reading frames in SEQ ID NO: 10084.

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FIGURE 123: Start codon analysis for SEO ID NO: 10084.

FIGURE 124: BLAST analysis of SEQ ID NO: 10210.

FIGURE 125: Epitope analysis of SEQ ID NO: 10210 by either (13A) Hopp & Woods or (13B) Kyte & Doolittle.

5 FIGURE 126: Reading frames in SEQ ID NO: 10299.

FIGURE 127: Reading frames in SEQ ID NO: 10505.

FIGURE 128: Reading frames in SEQ ID NO: 11563.

FIGURE 129: Reading frames in SEQ ID NO: 10033.

FIGURE 130: Alignment of SEQ ID NO: 9997 and SEQ ID NO: 10033.

10 FIGURE 131: Reading frames in SEQ ID NO: 10299.

FIGURE 132: Reading frames in SEQ ID NO: 10505...

FIGURE 133: Western Blot of SARS protease purification fractions.

FIGURE 134: Cleavage of DABCYL-EDANS (a fluorescent tagged peptide with a SARS protease cleavage site) by SARS protease at different concentrations. The graph shows

activity/concentration correlations with no protease (*), 0.95 uM protease (*) and 2.86 uM protease (*).

In the event of a discrepancy between a sequence in the sequence listing and a sequence in the drawings, the drawings should take precedence.

DETAILED DESCRIPTION OF THE INVENTION

20 The practice of the present invention will employ, unless otherwise indicated, conventional methods of chemistry, biochemistry, molecular biology, immunology and pharmacology, within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Remington's Pharmaceutical Sciences, Mack Publishing Company, Easton, Pa., 19th Edition (1995); Methods In Enzymology (S. Colowick and N. Kaplan, eds., Academic Press, Inc.); and Handbook of 25 Experimental Immunology, Vols. I-IV (D.M. Weir and C.C. Blackwell, eds., 1986, Blackwell Scientific Publications); Sambrook, et al., Molecular Cloning: A Laboratory Manual (2nd Edition, 1989); Handbook of Surface and Colloidal Chemistry (Birdi, K.S. ed., CRC Press, 1997); Short Protocols in Molecular Biology, 4th ed, (Ausubel et al. eds., 1999, John Wiley & Sons); Molecular Biology Techniques: An Intensive Laboratory Course, (Ream et al., eds., 1998, 30 Academic Press); PCR (Introduction to Biotechniques Series), 2nd ed. (Newton & Graham eds., 1997, Springer Verlag); Peters and Dalrymple, Fields Virology (2d ed), Fields et al. (eds.), B.N. Raven Press, New York, NY.

All publications, patents and patent applications cited herein, are hereby incorporated by reference in their entireties.

Severe Acute Respiratory Syndrome (SARS) virus has recently been identified as a new viral species. The SARS viral species includes the following isolates.

- two virus isolates described in Peiris et al. "Coronavirus as a possible cause of severe acute respiratory syndrome" Lancet published online at http://image.thelancet.com/extras/03ari3477web.pdf on April 8 2003, incorporated herein by reference in its entirety and the sequences deposited with GenBank at accession number AY268070.
- the isolates and viral sequences described in Drosten et al., "Identification of a Novel Coronavirus in Patients with Severe Acute Respiratory Syndrome", New England Journal of Medicine, published online at http://www.nejm.org on April 10, 2003.
 - the isolates and viral sequences described on the website of the WHO network on March 25 and 24, 2003.
- the isolates and viral sequences described in Tsang et al., "A Cluster of Cases of Severe
 Acute Respiratory Syndrome in Hong Kong" New England Journal of Medicine, published online at http://www.nejm.org on March 31, 2003.
 - the isolates and viral sequences described in Poutanen et al., "Identification of Severe Acute Respiratory Syndrome in Canada" New England Journal of Medicine, published online at http://www.nejm.org on March 31, 2003.

As described in the *Lancet* article, a 646 base pair polynucleotide from the SARS virus has weak homology to viruses of the family *Cornoaviridae*. The *Lancet* article further reports that a deduced amino acid sequence (of 215 amino acids) from this sequence has about 57% sequence homology to the RNA polymerase of bovine coronavirus and murine hepatitis virus.

25 Phylogenetic analysis of the protein sequences are also presented in the *Lancet* article showing that the polymerase sequence is most closely related to the group II coronaviruses.

Additional SARS viral isolates can be identified, isolated and/or sequenced by virologists skilled in the art. Virologists can readily identify new viral isolates as a SARS virus. Criteria which a virologist may use to identify new SARS isolates include: sequence homology of the new isolate to known SARS viral isolates; similar genomic organization of the new viral isolate to known SARS viral isolates; immunological (serologic) similarity or identity with known SARS viral isolates; pathology; and similarity of virion morphology with known SARS viral isolates; and similarity of infected cell morphology as that caused by known SARS viral isolates (visualized, for instance, by electron microscopy).

Methods for isolating and sequencing SARS viral isolates include the methods described by Peiris et al. in the Lancet paper. As reported in the Lancet paper, RNA from clinical samples

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can be reverse transcribed with random hexamers and cDNA can be amplified with primers having sequences of SEQ ID NOS: 6584 & 6585 in the presence of 2.5 mmol/L magnesium chloride (94°C for 1 min, 50°C for 1 min, and 72°C for 1 min).

Reverse transcription of a viral isolate using random hexamers can be accomplished in an RT-PCR assay as follows. Virus isolates are propagated on mammalian cells, particularly fetal rhesus kidney cells. Total RNA from virus-infected and virus-uninfected fetal rhesus kidney cells is then isolated. RNA samples are reverse transcribed with a primer having SEQ ID NO: 6586. cDNA can be amplified by a primer having SEQ ID NO: 6587. Unique PCR products (in size) in the infected cell preparation are then cloned and sequenced, and genetic homology of the sequence compared with those in GenBank.

One skilled in the art would be able to identify and clone additional genomic regions using a variety of standard cloning techniques, such as, for example, using random primer RT-PCR and detection of sequences overlapping one or more of the above sequences, and/or using oligonucleotide primers, e.g., degenerate primers, based on the sequences provided herein (see Figures 1-5, Figures 8-11, SEO ID NOS: 3-20).

Cloning, sequencing and identification of SARS virus by one skilled in the art can be further facilitated by the use of polynucleotide sequences, particularly RNA polymerase sequences, from related Coronaviruses.

Sequence homology of new viral isolates with the known SARS isolates described above can be readily determined by one skilled in the art. New SARS isolates may be identified by a percent homology of viral nucleotide sequences of 99%, 95%, 92%, 90%, 85%, or 80% homology of the new virus to known SARS viral polynucleotide sequences. New SARS isolates may also be identified by percent homology of 99%, 95%, 92%, 90%, 85%, or 80% homology of the polypeptides encoded by the polynucleotides of the new virus and the polypeptides encoded by known SARS virus.

New SARS isolates may also be identified by a percent homology of 99%, 95%, 92%, 90%, 85%, or 80% homology of the polynucleotide sequence for specific genomic regions for the new virus with the polynucleotide sequence for specific genomic regions of the known SARS viruses. Additionally, new SARS isolates may be identified by a percent homology of 99%, 95%, 92%, 90%, 85%, or 80% homology of the polypeptide sequence encoded by the polynucleotide of specific genomic regions of the new SARS virus to the polypeptide sequence encoded by the polynucleotides of specific regions of the known SARS virus. These genomic regions may include regions (e.g., gene products) which are typically in common among numerous coronaviruses, as well as group specific regions (e.g., antigenic groups), such as, for example, any one of the following genomic regions which could be readily identified by a virologist skilled in the art: 5'untranslated region (UTR), leader sequence, ORF1a, ORF1b,

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nonstructural protein 2 (NS2), hemagglutinin-esterase glycoprotein (HE) (also referred to as E3), spike glycoprotein (S) (also referred to as E2), ORF3a, ORF3b, ORF3x, nonstructural protein 4 (NS4), envelope (small membrane) protein (E) (also referred to as sM), membrane glycoprotein (M) (also referred to as E1), ORF5a, ORF5b, nucleocapsid phosphoprotein (N), ORF7a, ORF7b, intergenic sequences, 3'UTR, or RNA dependent RNA polymerase (pol). The SARS virus may have identifiable genomic regions with one or more the above-identified genomic regions. A SARS viral antigen includes a protein encoded by any one of these genomic regions. A SARS viral antigen may be a protein or a fragment thereof, which is highly conserved with coronaviruses. A SARS viral antigen may be a protein or fragment thereof, which is specific to the SARS virus (as compared to known cornaviruses). (See, Figures 1-5, Figures 8-11, SEQ ID NOS: 3-20).

One skilled in the art could also recognize electron microscopy of a SARS virus infected mammalian cell. Electron microscopy of SARS infected cells are shown in the *Lancet* paper. As discussed in the paper, electron microscopy of negative stained (3% potassium phosphotungstate, pH 7.0) ultracentrifuged cell-culture extracts of SARS infected fetal rhesus kidney cells show the presence of pleomorphic enveloped virus particles of around 80-90 nm (range 70-130 nm) in diameter with surface morphology compatible with a coronavirus (see *Lancet* paper, Figure 1). Thin-section electron microscopy of infected cells reveals virus particles of 55-90 nm diameter within smooth walled vesicles in the cytoplasm (see *Lancet* paper, Figure 2B). Electron microscopy can also be used to observe virus particles at the cell surface. Electron microscopy of a human lung biopsy sample depicts similar viral morphology. See *Lancet* paper Figure 2A.

I. SARS POLYPEPTIDES AND POLYNUCLEOTIDES

The invention relates to nucleic acids and proteins from SARS virus. Such polynucleotides and polypeptides are exemplified further below.

In one embodiment, the polynucleotides of the invention do not include one of the following five primers, disclosed at http://content.nejm.org/cgi/reprint/NEJMoa030781v2.pdf: SEQ ID NOS: 6034-38.

The invention also includes polynucleotide sequences which can be used as probes for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in SEQ ID NOS: 21-1020. The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in SEQ ID NOS: 21-1020.

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The invention includes a polypeptide sequence comprising an amino acid sequence from the sequence shown in Figure 23. Such amino acid sequences are SEQ ID NOS: 6588-6809. The invention includes polypeptides comprising an amino acid sequence having sequence identity to these sequences, and the invention includes a fragment of a polypeptide comprising one of these sequences.

The invention includes a polypeptide comprising an amino acid sequence from the sequence shown in Figure 24. Such amino acid sequences are SEQ ID NOS: 6810-7179. The invention includes a protein comprising an amino acid sequence having sequence identity to these sequences, and the invention includes a fragment of a protein comprising one of these sequences.

The invention includes a protein comprising SEQ ID NO: 6039. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 6039. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 6039. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6039, or a fragment thereof. The invention includes a diagnostic kit comprising a polynucleotide sequence encoding SEQ ID NO: 6039, or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising SEQ ID NO: 6039, or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 6039, or a fragment thereof. SEQ ID NO: 6039 demonstrates functional homology with ORF1a of coronaviruses.

Predicted transmembrane or hydrophobic regions of SEQ ID NO: 6039 are identified below. Although the polyprotein of coronaviruses is proteolytically cleaved into numerous smaller proteins, hydrophobic domains in the polyprotein are known to mediate the membrane association of the replication complex and to be able to dramatically alter the architecture of host cell membranes. Accordingly, the hydrophobic domains of the polyprotein are targets for genetic mutation to develop attenuated SARS virus vaccines. The hydrophobic domains are also targets for small molecule inhibitors of the SARS virus. The hydrophobic domains may also be used to generate antibodies specific to those regions to treat or prevent SARS virus infection.

Predicted Transmembrane Helices in SEQ ID NO: 6039

Inside to outside helices: 43 found

The sequence positions in brackets denominate the core region. Only scores above 500 are considered significant.

			from			to	score	center			from			to	score	center		
	100	(100)	118	(116)	103	107	94	(97)	118	(112)	291	104		
	473	(473)	488	(488)	1003	481	400	i	400)	418	i	415)	243	407		
	529	Ċ	532)	549	i	549)	541	539	473	i	473)	488	i	488)	1113	481		
	584	Ċ	584)	606	ĺ	601)	1049	594	523	i	528)	548	i	548)	285	538		
	773	i	7731	701	i	7801	51/	782	503	i	5831	606	ì	6011	662	503		

Outside to inside helices :

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1071	(1071) 1089	(1086)	243	1078	776	(776) 791	(791)	1435	783
1121	(1121)1137	(1137)	459	1130	1068	(1071)1089	(1086)	370	1078
1679	(1679)1696	(1696)	404	1686	1121	(1121)1137	(1137)	455	1130
2098	(2102)2119	(2116)	509	2109	1679	(1679)1696	(1694)	340	1686
2145	(2145)2160	(2160)	797	2153	2098	(2098)2119	(2116)	678	2109
2206	(2209)2224	(2224)	2686	2216	2148	(2148)2163	(2163)	434	2155
2316	(2316) 2332	(2332)	2123	2325	2208	(2210)2231	(2226)	2389	2219
2335	(2339)2358	(2354)	2101	2346	2309	(2309)2332	(2326)	1773	2318
2373	(2373)2390	(2390)	532	2380	2342	(2342)2368	(2360)	1666	2353
2597	(2600)2615	(2615)	307	2607	2373	(2373)2390	(2390)	254	2380
2753	(2753)2770	(2768)	2242	2760	2753	(2755)2770	(2770)	2119	2763
2831	(2833)2854	(2851)	759	2841	2832	(2835) 2854	(2851)	687	. 2844
2879	(2882) 2900	(2897)	526	2889	2858	(2858) 2873	(2873)	253	2866
2990	(2996)3012	(3010)	1289	3003	2879	(2882)2899	(2899)	400	2889
3024	(3024)3042	(3039)	2281	3032	2990	(2990)3005	(3005)	875	2998
3054	(3058) 3075	(3072)	2536	3065	. 3020	(3024)3042	(3042)	2795	3032
3105	(3109)3127	(3123)	2010	3116	3059	(3059)3075	(3075)	2137	3067
3143	(3143)3163	(3159)	184	3152	3105	(3108)3127	(3123)	1902	3115
3253	(3255) 3272	(3272)	319	3262	3142	(3145)3162	(3162)	540	3152
3346	(3346)3366	(3366)	203	3356	3343	(3351)3366	(3366)	496	3358
3375	(3375)3392	(3392)	305	3384	3437	(3437)3453	(3453)	848	3444
3438	(3438)3455	(3453)	1021	3445	3489	(3491)3508	(3505)	302	3498
	(3567)3584	(3581)		3574	3560	(3560)3577	(3577)	1460	3569
3589	(3589)3606	(3604)	2018	3596	3591	(3591)3606	(3606)	2193	3598
		(3629)	2304	3621	3610	(3610)3627	(3627)		3620
3659	(3659)3674	(3674)	1561	3667	3656	(3658)3678	(3675)	1240	3668
3756	(3758) 3777	(3774)	2352	3767	3681	(3684)3701	(3699)	590	3691
3890	(3890)3904	(3904)	485	3897	3710	(3713)3738	(3728)	1696	3721
3916	(3919)3934	(3934)	241	3926	3723	(3723)3738	(3738)	1670	3730
4035	(4035) 4051	(4051)	335	4044		(3760)3777	(3775)	2367	3767
4217	(4217) 4232	(4232)	272	4224	3881	(3884)3902	(3900)	249	3892
4239	(4239) 4257	(4254)	402	4247	4099	(4099)4114	(4114)	389	4106
					4234	(4234) 4254	(4249)	325	4241
					4338	(4341)4360	(4360)	505	4348

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6039, wherein said fragment comprises an amino acid sequence including one or more of the hydrophobic transmembrane sequences identified above. The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6039 wherein said fragment comprises one or more of the following polypeptide sequences of SEQ ID NO: 6039: 473-488, 529-549, 584-606, 773-791, 2098-2119, 2145-2160, 2206-2224, 2316-2332, 2335-2358, 2373-2390, 2753-2770, 2831-2854, 2879-2900, 2990-3012, 3024-3042, 3054-3075, 3105-3127, 3438-3455, 3559-3584, 3589-3606, 3611-3629, 3659-3674, 3756-3777, 473-488, 583-606, 776-791, 2098-2119, 2208-2231, 2309-2332, 2342-2368, 2753-2770, 2832-2854, 2990-3005, 3020-3042, 3059-3075, 3105-3127, 3142-3162, 3437-3453, 3560-3577, 3591-3606, 3610-3627, 3656-3678, 3710-3738, 3723-3738, and 3760-3777. Preferably, the fragment comprises one or more of the following polypeptide sequences of SEQ ID NO: 6039: 2206-2224, 2316-2332, 2335-2358, 2753-2770, 3024-3042, 3054-3075, 3105-3127, 3589-3606, 3611-3629, 3756-3777, 2208-2231, 2753-2770, 3020-3042, 3059-3075, and 3591-3606. Preferably, the fragment comprises one or more of the following

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polypeptide sequences of SEQ ID NO: 6039: 2206-2224 and 3020-3042. The invention also includes polynucleotides encoding each of the polypeptide fragments identified above.

The invention includes an attenuated SARS virus wherein said attenuated SARS virus contains an addition, deletion or substitution in the polynucleotides encoding for one of the hydrophobic domains identified above. The invention also includes a method for creating an attenuated SARS virus comprising mutating a SARS virus by adding, deleting or substituting the viral genome of the SARS virus to alter the coding of one or more of the hydrophobic domains of SEQ ID NO: 6039 identified above.

The invention includes an antibody which specifically identifies one or more of the hydrophobic regions of SEQ ID NO: 6039 identified above. The invention includes a small molecule which binds to, interferes with the hydrophobicity of or otherwise disrupts one or more of the hydrophobic regions of SEQ ID NO: 6039 identified above.

Predicted N-glycosylation sites of SEQ ID NO: 6039 are identified in the chart below. Prediction of N-glycosylation sites in SEQ ID NO: 6039

15	IICUICCION OI	. GIYCOSY	TACTOR	SICOS	III SEQ II	3 NO: 6039			
	Pos	sition			Potential	Jury	NGlyc		
						agreement	result		
	48	NGTC SEQ	ID NO:	7180	0.6371	(7/9)	+		
	.389	NHSN SEQ	ID NO:	7181	0.6132	(6/9)	+		
20	916	NFSS SEQ	ID NO:	7182	0.5807	(7/9)	+		
	1628	NHTK SEQ	ID NO:	7183	0.5610	(7/9)	+		
	1696	NKTV SEQ	ID NO:	7184	0.5297	(5/9)	+		
	2031	NPTI SEQ	ID NO:	9764	0.5299	(5/9)	+ .	WARNING:	PRO-
	X1.					,,			
25	2249	NSSN SEQ	ID NO:	7185	0.6329	(9/9)	++		
	2459	NPTD SEQ	ID NO:	9765	0.5599	(6/9)	+	WARNING:	PRO-
	X1.								
	2685	NVSL SEQ	ID NO:	7186	0.6071	(8/9)	+		
		NATE SEO				(7/9)	+		
30		_							

Accordingly, the invention comprises a fragment of SEQ ID NO: 6039 wherein said fragment comprises an amino acid sequence which includes one or more of the N-glycosylation sites identified above. Preferably, the fragment comprises one or more sequences selected from the group consisting of SEQ ID NOS: 7180-7187 & 9764-9765. Preferably, the fragment comprises the amino acid sequence NSSN (SEQ ID NO: 7185).

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6039 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polypucleotide encoding such a polypeptide.

T-epitopes for SEQ ID NO: 6039 are identified in Table 13. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified as SEQ ID NOS: 7400-7639; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the

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polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified as SEQ ID NOS: 7400-7639, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus.

The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The ORF1a and ORF1b sequences of coronaviruses are typically translated as a single ORF1ab polyprotein. Slippage of the ribosome during translation generates an a-1 frameshift. One region of such slippage is illustrated below:

Accordingly, the invention includes a polypeptide comprising SEQ ID NO: 7232. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 7232. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 7232. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 7232 or a fragment thereof. The invention includes a diagnostic kit comprising a polypucleotide sequence encoding SEQ ID NO: 7232 or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising SEQ ID NO: 7232 or a

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fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 7232 or a fragment thereof.

The invention also includes a polypeptide comprising amino acid sequence X_1 - X_2 - X_3 , where X_1 is SEQ ID NO: 7233, X_2 is from one to ten amino acids, and X_3 is SEQ ID NO: 7234. X_2 can comprise any sequence of one to ten amino acids (SEQ ID NOS: 7235-7244) but, in preferred embodiments, X_2 is selected from the group consisting of F, FL, FLN, FLNR (SEQ ID NO: 7245), FLNRV (SEQ ID NO: 7246) and FLNRVC (SEQ ID NO: 7247). Preferably, X_2 is SEO ID NO: 7247. These preferred embodiments are shown as SEO ID NOS: 7248-7253.

The invention includes a polypeptide comprising an amino acid sequence having sequence identity to said amino acid sequences X_1 - X_2 - X_3 . The invention includes a fragment of a polypeptide comprising said amino acid sequences X_1 - X_2 - X_3 . The invention includes a diagnostic kit comprising a polypeptide comprising said amino acid sequences X_1 - X_2 - X_3 or a fragment thereof. The invention includes a diagnostic kit comprising a polypucleotide sequence encoding said amino acid sequences X_1 - X_2 - X_3 or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising said amino acid sequences X_1 - X_2 - X_3 or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising said amino acid sequences X_1 - X_2 - X_3 or a fragment thereof.

The amino acid sequences X_1 - X_2 - X_3 (i.e. SEQ ID NOS: 7235-7244) demonstrate functional homology with the polyprotein of murine hepatitis virus. This polyprotein is cleaved to produce multiple proteins. Proteins which can be generated from the X_1 - X_2 - X_3 polyprotein, where X_2 is six amino acids (SEQ ID NO: 7240) are listed below.

Mouse virus protein	Coordinates in Mouse virus	Coordinates in SEQ ID NO: 7240
Nsp2	3334-3636	3241-3546
Nsp3	3637-3923	3547-3836
Nsp4	3924-4015 (or 4012)	3837-3919
Nsp5	4016 (or 4013)-4209	3920-4117
Nsp6	4210-4319	4118-4230
Nsp7	4320-4456	4231-4369
Nsp9	4457-5384	4370-5301
Nsp10	5385-5984	5302-5902
Nsp11	5985-6505	5903-6429
Nsp12	6506-6879	6430-6775
Nsp13	6880-7178	6776-7073

The invention includes a fragment of the amino acid sequence $X_1-X_2-X_3$ (i.e. SEQ ID NOS: 7235-7244) wherein the fragment comprises one of the polypeptide sequences identified in the above table. The invention further includes a fragment of the amino acid sequence $X_1-X_2-X_3$ wherein said fragment comprises a polypeptide sequence which has a serine at its N-terminus and a glutamine at its C-terminus. The invention further includes a fragment of the amino acid sequence $X_1-X_2-X_3$ wherein said fragment comprises a polypeptide sequence which has an

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Alanine at its N-terminus and a glutamine at its C-terminus. The invention further includes a fragment of the amino acid sequence X_1 - X_2 - X_3 wherein said fragment comprises a polypeptide sequence which has a Asparagine at its N-terminus and a glutamine at its C-terminus. The invention further includes a fragment of the amino acid sequence X_1 - X_2 - X_3 wherein said fragment comprises a Cysteine at its N-terminus and a Glutamine at its C-terminus. Each of the fragments identified above can be used in fusion proteins.

The invention includes a diagnostic kit comprising a polypeptide comprising at least one of the fragments of the amino acid sequence X_1 - X_2 - X_3 (i.e. SEQ ID NOS: 7235-7244) identified in the above paragraph. The invention includes a diagnostic kit comprising a polynucleotide sequence encoding at least one of the fragments of the amino acid sequence X_1 - X_2 - X_3 identified in the above paragraph. The invention includes an immunogenic composition comprising a polypeptide comprising at least one of the fragments of the amino acid sequence X_1 - X_2 - X_3 identified in the above paragraph. The invention includes an antibody which recognizes a polypeptide comprising at least one of the fragments of the amino acid sequence X_1 - X_2 - X_3 identified in the above paragraph.

Predicted N-glycosylation sites of the amino acid sequence X_1 - X_2 - X_3 when X_2 is six amino acids are identified at the asparagines located at the following amino acid positions 48; 389; 556; 916; 1628; 1696; 1899; 2079; 2249; 2252; 2507; 2685; 3303; 3373; 3382; 3720; 4150; 4233; 4240; 5016; 5280; 5403; 5558; 5650; 5905; 6031; 6130; 6474; 6918; 6973. Accordingly, the invention comprises a fragment of SEQ ID NO: 7239 wherein said fragment is at least ten amino acids and wherein said fragment comprises one or more of the asparagines from the amino acid positions of SEQ ID NO: 7239 selected from the group consisting of 8; 389; 556; 916; 1628; 1696; 1899; 2079; 2249; 2252; 2507; 2685; 3303; 3373; 3382; 3720; 4150; 4233; 4240; 5016; 5280; 5403; 5558; 5650; 5905; 6031; 6130; 6474; 6918; and 6973.

A zinc binding region 2 site within SEQ ID NOS: 7235-7244 is identified at amino acid residues 2102-2112 (SEQ ID NO: 7254 HGIAAINSVFW). The polypeptide of SEQ ID NOS: 7235-7244 will be processed by the SARS virus into multiple peptides. This zinc binding region falls within the nsp1 region of the polypeptide. SEQ ID NO: 7254 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 7254. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 7254. The invention includes a method of screening SEQ ID NO: 7254 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 7254 in a host cell. The invention includes a fragment of SEQ ID NOS: 7235-7244, wherein said fragment comprises SEQ ID NO: 7254. The invention includes a polypeptide comprising SEQ ID NO: 7254 wherein said polypeptide is complexed with a zinc ion. The invention includes a small molecule which

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prevents a zinc ion from complexing with the polypeptide of SEQ ID NO: 7254. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 7254.

The polyprotein encoded by the SARS virus will contain at least two protease domains: a papain-like cystein protease (PLP) and a chymotrypsin-picornavirus 3C-like protease (3CLp). (There may be more than one copy of the PLP domain). These proteases function to cleave the polyprotein into multiple smaller proteins. The 3C-like protease, also known as the "main protease" or Mpro, is itself cleaved from the polyprotein by its own autoprotease activity. See generally, Chapter 35 of Fields Virology (2nd ed), Fields et al. (eds.), B.N. Raven Press, New York, NY, and Anand et al., EMBO Journal (2002) 21 (13): 3213-3224. This 3CLp generally corresponds with the Nsp2 region identified above.

The SARS virus 3CLp protein is further characterized by SEQ ID NO: 6569 (also SEQ ID NO: 9769), as shown in FIGURE 15.

FIGURE 16 also illustrates the SARS virus 3CLp, in allignment with the 3CLp of avian infectious bronchitis (IBV; SEQ ID NO: 6570), mouse hepatitis virus (MHV; SEQ ID NO: 6571), and bovine coronavirus (BCoV; SEQ ID NO: 6572). Accordingly, the invention includes a polypeptide sequence comprising SEQ ID NO: 6569, or a fragment thereof, or a polypeptide sequence having sequence identity thereto. The invention further includes a polynucleotide sequence encoding SEQ ID NO: 6569, or a fragment thereof. The invention includes a polynucleotide sequence encoding a polypeptide sequence having sequence identity to SEQ ID NO: 6569.

The invention further includes a method of screening for an inhibitor of the SARS virus 3CLp protein. In one embodiment, the invention includes a method of screening for an inhibitor of SEQ ID NO: 6569. The invention includes a method of recombinantly expressing the SARS virus 3CLp protein in a host cell. The invention includes a method of recombinantly expressing a polypeptide sequence comprising SEQ ID NO: 6569 or an enzymatically active fragment thereof or a polypeptide sequence having sequence identity thereto. The invention includes a small molecule which inhibits or reduces the proteolytic activity of the SARS virus 3CLp protein. The invention includes a small molecule which inhibits or reduced the proteolytic activity of the polypeptide comprising SEQ ID NO: 6569.

Catalytic residues of the SARS virus 3CLp are identified in FIGURE 15 and 16.

Specifically, a catalytic histidine and a catalytic cysteine are identified. Such catalytic sites are targets for small molecules which could inhibit or reduce the protease activity of 3CLp.

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6569, wherein said fragment comprises at least one catalytic site. Preferably, the catalytic site is selected from the group consisting of the indicated catalytic histidine and the catalytic cysteine in FIGURE 15 and 16. The invention includes a polypucleotide encoding a polypeptide, wherein

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said polypeptide comprises a fragment of SEQ ID NO: 6569, wherein said fragment comprises at least one catalytic site. Preferably, the catalytic site is selected from the group consisting of the indicated catalytic histidine and the catalytic cysteine.

The invention further includes a method of screening a compound library to identify a small molecule which inhbits a catalytic site of a SARS virus 3CLp. Preferably, the 3CLp comprises SEQ ID NO: 6569, or a fragment thereof, or a sequence having sequence identity thereto. The catalytic site is preferably selected from the group consisting of the indicated catalytic histidine and the catalytic cysteine in FIGURE 15 and 16

The invention includes a small molecule which inhibits the catalytic site of a SARS virus 3CLp. Preferably, the 3CLp comprises SEQ ID NO: 6569, or a fragment thereof, or a sequence having sequence identity thereto. The catalytic site is preferably selected from the group consisting of the indicated catalytic histidine and the catalytic cysteine in FIGURE 15 and 16.

Residues of the substrate site of the SARS virus 3CLp are identified in FIGURE 15 and 16. Specifically, a substrate site is indicated at a phenylalanine, a tyrosine and a histidine. Such substrate sites are targets for small molecules which could inhibit or reduce the protease activity of 3CLp. Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6569, wherein said fragment comprises at least one substrate site. Preferably, the substrate site is selected from the group consisting of the indicated substrate phenylalanine, tyrosine and histidine in FIGURE 15 and 16. The invention includes a polynucleotide encoding a polypeptide, wherein said polypeptide comprises a fragment of SEQ ID NO: 6569, wherein said fragment comprises at least one substrate site. Preferably, the substrate site is selected from the group consisting of the indicated substrate phenylalanine, tyrosine and histidine in FIGURE 15 and 16.

The invention further includes a method of screening a compound library to identify a small molecule which blocks a substrate site of a SARS virus 3CLp. Preferably, the 3CLp comprises SEQ ID NO: 6569, or a fragment thereof, or a sequence having sequence identity thereto. The substrate site is preferably selected from the group consisting of the indicated substrate phenylalanine, tyrosine and histidine in FIGURE 15 and 16.

The invention includes a small molecule which inhibits the substrate site of a SARS virus 3CLp. Preferably, the 3CLp comprises SEQ ID NO: 6569, or a fragment thereof, or a sequence having sequence identity thereto. The substrate site is preferably selected from the group consisting of the indicated substrate phenylalanine, tyrosine and histidine in FIGURE 15 and 16.

The invention further includes a diagnostic kit comprising a polynucleotide encoding a SARS virus 3CLp or a fragment thereof. Preferably, the SARS virus 3CLp comprising SEQ ID NO: 6569 or a fragment thereof or a polypeptide sequence having sequence identity thereto. Preferably, the fragment comprising one or more sites selected from the group consisting of a

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catalytic site and a substrate site. Preferably, the catalytic site is selected from the group consisting of one or more of the sites identified in FIGURE 15 and 16. Preferably, the substrate site is selected from the group consisting of one or more of the sites identified in FIGURE 15 and 16.

The invention further comprises a diagnostic kit comprising an antibody specific to a SARS virus 3CLp or a fragment thereof. Preferably, the antibody is specific to the polypeptide comprising SEQ ID NO: 6569 or a fragment thereof or a polypeptide sequence having sequence identity thereto. Preferably, the antibody is specific to one or more sites of a SARS virus 3CLp selected from the group consisting of a catalytic site and a substrate site. Preferably, the catalytic site is selected from the group consisting of one or more of the sites identified in FIGURE 15 and 16. Preferably, the substrate site is selected from the group consisting of one or more of the sites identified in FIGURE 15 and 16.

The invention includes a polypeptide comprising an amino acid sequence from the sequence shown in Figure 25. The two amino acid sequences within Figure 25, separated by a *, are SEQ ID NOS: 7188 & 7189. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to the figure 25 translation. The invention includes a fragment of a polypeptide comprising the figure 25 sequence. The invention includes a diagnostic kit comprising a polypeptide comprising the figure 25 translation, or a fragment thereof. The invention includes a diagnostic kit comprising a polypucleotide sequence encoding the figure 25 translation, or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising the figure 25 translation, or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising the figure 25 sequence, or a fragment thereof. The figure 25 sequence demonstrates functional homology with ORF1b of coronaviruses.

SEQ ID NO: 7188 is an open reading frame within Figure 25. The invention includes a polypeptide comprising SEQ ID NO: 7188. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 7188. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 7188. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 7188, or a fragment thereof. The invention includes a diagnostic kit comprising a polypeptide sequence encoding SEQ ID NO: 7188, or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising SEQ ID NO: 7188, or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 7188, or a fragment thereof.

SEQ ID NO: 7190 is an open reading frame within SEQ ID NO: 7188. The invention includes a polypeptide comprising SEQ ID NO: 7190, a fragment thereof or a polypeptide having sequence identity thereto. The invention further includes a polynucleotide encoding SEQ

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ID NO: 7190, a fragment thereof or a polypeptide sequence having sequence identity thereto. An example of a polynucleotide encoding SEO ID NO: 7190 is given as SEO ID NO: 7191.

SEQ ID NO: 7188 also contains an open reading frame comprising SEQ ID NO: 6042. The invention includes a polypeptide comprising SEQ ID NO: 6042. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 6042. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 6042. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6042, or a fragment thereof. The invention includes a diagnostic kit comprising a polypucleotide sequence encoding SEQ ID NO: 6042, or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising SEQ ID NO: 6042, or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 6042, or a fragment thereof. SEQ ID NO: 6042 demonstrates functional homology to a coronavirus spike protein.

Predicted transmembrane regions of SEQ ID NO: 6042 are identified below.

Predicted Transmembrane helices of SEQ ID NO: 6042

The sequence positions in brackets denominate the core region. Only scores above 500 are considered significant.

Insid	le	to o	utsio	ie	helic	es	:	18	found	(Outsi	de	to.	insi	de	helice	s:	1	found	
	3	from			to	sc	ore	. c	enter			Í	rom			to	scor	e i	center	
1	(1)	16	(16)		959	•	9		1	(1)	17	(17)	68	4	10	
233	(237)	257	(252)		905	,	244		222	(222)	240	(237)	23	8	229	
345	(347)	364	(361)		490	1	354		244	(247)	264	(264)	61	.3	254	
345	(354)	369	(369)		420	1	362		349	Ĺ	355)	369	i	369)	31	4	362	
497	(497)	513	(513)		239	,	506		496	Ċ	496)	511	i	511)	48	8	503	
573	(573)	588	(588)		811		580		573	Ĺ	573)	591	i	591)	71	2	581	
645	(648)	666	(663)		302		656		650	i	652)	666	i	666)	47	4	659	
690	(696)	714	(711)		428	:	704							696)	19	0	686	
857	Ĺ	860)	882	i	874)	1	508	:	867							711)	21	-	704	
1031	(3	1031)	1046	(:	1046)		446		1039							886)	117	_	876	
1199	Ü	1203)	1219	Ċ	1217)	2	667		1210							1215)	322		1200	

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SEQ ID NO: 6042, the spike protein, is a surface exposed polypeptide. Recombinant expression of a protein can be hindered by hydrophobic transmembrane regions. Accordingly, the invention includes a polypeptide comprising SEQ ID NO: 6042 wherein one or more of the hydrophobic regions identified above is removed. The invention further includes a polypucleotide encoding such a polypeptide. The invention includes recombinantly expressing the protein in a host cell. Primers for amplifying the gene for spike protein and fragments thereof, such as fragments encoding the soluble ectodomain, include SEQ ID NOS: 9753-9763 (Xiao et al. (2003) Biochem Biophys Res Comm 312:1159-1164).

Further characterization of SEQ ID NO: 6042 is set forth below.

PSORT --- Prediction of Protein Localization Sites

```
version 6.4(WWW)
       SEQ ID NO: 6042 - 1255 Residues
       Species classification: 4
5
      *** Reasoning Step: 1
      Preliminary Calculation of ALOM (threshold: 0.5)
            count: 2
            Position of the most N-terminal TMS: 496 at i=2
10
      MTOP: membrane topology (Hartmann et al.)
            T(middle): 503 Charge diffirence(C-N): 1.0
      McG: Examining signal sequence (McGeoch)
            Length of UR: 13
            Peak Value of UR: 3.28
           Net Charge of CR: 0
15
                                    8.66
            Discriminant Score:
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): 5.94
            Possible cleavage site: 13
       >>> Seems to have a cleavable N-term signal seq.
20
       Amino Acid Composition of Predicted Mature Form:
          calculated from 14
       ALOM new cnt: 1 ** thrshld changed to -2
       Cleavable signal was detected in ALOM?: OB
       ALOM: finding transmembrane regions (Klein et al.)
25
            count: 1 value: -12.26 threshold: -2.0
                       Likelihood =-12.26 Transmembrane 1202-1218 (1194-1228)
            INTEGRAL
            PERIPHERAL Likelihood = 0.16
            modified ALOM score: 2.55
       >>> Seems to be a Type Ia membrane protein
30
            The cytoplasmic tail is from 1219 to 1255 (37 Residues)
       Rule: vesicular pathway
       Rule: vesicular pathway
       Rule: vesicular pathway
35
       (14) or uncleavable?
       Gavel: Examining the boundary of mitochondrial targeting seq.
             motif at: 14
            Uncleavable? Ipos set to: 24
       Discrimination of mitochondrial target seq.:
40
            positive (2.18)
       Rule: vesicular pathway
       Rule: vesicular pathway
       Rule: vesicular pathway
45
       *** Reasoning Step: 2
       KDEL
              Count: 0
       Checking apolar signal for intramitochondrial sorting
         (Gavel position 24) from: 1 to: 10 Score: 8.0
        SKL motif (signal for peroxisomal protein):
50
            pos: 964(1255), count: 1 SRL
            SKL score (peroxisome): 0.1
        Amino Acid Composition Tendency for Peroxisome:
            AAC not from the N-term., score modified
55
        Peroxisomal proteins? Status: notclr
            AAC score (peroxisome): 0.079
        Amino Acid Composition tendency for lysosomal proteins
            score: 0.39 Status: notclr
        GY motif in the tail of typeIa? (lysosomal)
        Checking the amount of Basic Residues (nucleus)
 60
        Checking the 4 residue pattern for Nuclear Targeting
```

```
Checking the 7 residue pattern for Nuclear Targeting Checking the Robbins & Dingwall consensus (nucleus) Checking the RNA binding motif (nucleus or cytoplasm) Nuclear Signal Status: negative (0.00)

Type Ia is favored for plasma memb. proteins Checking the NPXY motif..
Checking the YXFF motif..
Checking N-myristoylation..
```

10 ---- Final Results ---plasma membrane --- Certainty= 0.460(Affirmative) < succ>
microbody (peroxisome) --- Certainty= 0.171(Affirmative) < succ>
endoplasmic reticulum (membrane) --- Certainty= 0.100(Affirmative) < succ>
endoplasmic reticulum (lumen) --- Certainty= 0.100(Affirmative) < succ>

SEQ ID NO: 6042 appears to have a N-terminus signaling region, followed by a surface exposed region, followed by a transmembrane region followed by a C-terminus cytoplasmic domain region. Accordingly, the invention includes an immunogenic, surface exposed fragment of SEQ ID NO: 6042. Preferably, said fragment comprises an amino acid sequence which does not include the last 50 amino acids of the C-terminus of SEQ ID NO: 6042. Preferably, said fragment comprises an amino acid sequence which does not include the last 70 amino acids of the C-terminus of SEQ ID NO: 6042. Preferably, said fragment does not include a transdomain region of SEQ ID NO: 6042. Preferably, said fragment does not include a C-terminus cytoplasmic domain of SEQ ID NO: 6042. Preferably, said fragment does not include a Nterminus signal sequence. Preferably, said fragment does not include amino acids 1-10 of the Nterminus of SEQ ID NO: 6042. Preferably, said fragment does not include amino acids 1-14 of the N-terminus of SEQ ID NO: 6042. Two oligopeptide fragments of SEQ ID NO: 6042 that are able to elicit anti-spike antibodies are SEQ ID NOS: 7398 & 7399, as described (with additional C-terminus cysteines) by Xiao et al. (2003) Biochem Biophys Res Comm 312:1159-1164. C-terminal truncations of spike protein, with removal of part of the cytoplasmic region, or removal upto and including the transmembrane region, are described by Yang et al. (2004)

A variant of SEQ ID NO: 6042 that is included within the invention is SEQ ID NO: 9962. Compared to SEQ ID NO: 6042, this sequence has Ser at residue 581 instead of Ala, and has Phe at residue 1152 instead of Leu.

The spike protein of coronaviruses may be cleaved into two separate chains into S1 and S2. The chains may remain associated together to form a dimer or a trimer. Accordingly, the invention includes a polypeptide comprising SEQ ID NO: 6042 wherein said polypeptide has been cleaved into S1 and S2 domains. The invention further includes a polypeptide comprising SEQ ID NO: 6042 wherein amino acids 1-10, preferably amino acids 1-14 of the N-terminus are removed and further wherein SEQ ID NO: 6042 is cleaved into S1 and S2 domains. Preferably the polypeptide is in the form of a trimer.

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Nature 428:561-564.

The spike protein appears to form an alpha-helical structure in the transmembrane region of the protein, preferably in the S2 domain. This alpha-helical structure is thought to associate with at least two additional spike proteins to form a trimer. Helical or coiled regions of the spike protein are identified below. Predicted coiled-coils of SEQ ID NO: 6042 (spike protein) are at amino acids 900-1005 and 1151-1185 (see Figure 12).

Accordingly, the invention comprises a polypeptide sequence comprising a fragment of SEQ ID NO: 6042 wherein said fragment includes a coiled region of SEQ ID NO: 6042. Said fragment preferably includes the amino acid sequences selected from the group consisting of amino acid positions 900 to 1005 and amino acid positions 1151 to 1185 of SEQ ID NO: 6042. The invention comprises a polypeptide sequence comprising a fragment of SEQ ID NO: 6042, wherein said fragment does not include a coiled region of SEQ ID NO: 6042. Said fragment preferably includes the amino acid sequences selected from the group consisting of amino acid positions 900 to 1005 and amino acid positions 1151 and 1185 of SEO ID NO: 6042.

The spike protein is believed to play an integral role in fusion and infection of Coronaviruses with mammalian host cells. Analysis of coronavirus spike proteins as well as similar surface proteins in other viruses has identified at least two structural motifs, typically located within the S2 domain, associated with this fusion event: heptad repeats (HR) and membrane fusion peptides.

At least two 4,3 hydrophobic heptad repeat (HR) domains are typically found in the ectodomain of the S2 domain of Coronaviruses. One heptad repeat region (HR1) is typically located adjacent to a fusion peptide while a second heptad region (HR2) is typically located near the C-terminus of the S2 domain, close to the transmembrane anchor. Heptad repeats are characteristic of coiled-coil structures and the heptad repeats found in viral surface proteins (such as coronavirus spike protein) are thought to form bundled helix structures which are involved in viral entry. See Bosch et al., J. Virology (2003) 77:8801-8811 (Figure 1B of this reference illustrates an alignment of the HR1 and HR2 regions of five coronaviruses along with SARS, annotated "HCov-SARS").

Heptad repeats generally contain a repeating structure of seven amino acids, designated a-b-c-d-e-f-g, where hydrophobic sidechains of residues a and d typically form an apolar stripe, and electrostatic interactions are found in residues e and g. Position a is most frequently Leu, Ile or Ala and position d is usually Leu or Ala. Residues e and g are often Glu or Gln, with Arg and Lys also prominent at position g. Charged residues are common to positions g, g and g as these residues may be in contact with solvent. Exceptions to these general parameters are known. For instance Pro residues are sometimes found within the heptad.

The HR1 and HR2 sequences of an MHV strain have been postulated to assemble into a thermostable, oligomeric, alphahelical rold-like complex, with the HR1 and HR2 helices

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oriented in an antiparallel manner. *Id.* In this same study, HR2 was asserted to be a strong inhitibor of both virus entry into the cell and cell-cell fusion.

HR1 and HR2 sequences have been identified in the SARS virus genome. The SARS virus HR1 region comprises approximately amino acids 879 to 1005 of SEQ ID NO: 6042 or fragments thereof capable of forming at least one alpha-helical turn. Preferably, said fragments comprise at least 7 (e.g., at least 14, 21, 28, 35, 42, 49 or 56) amino acid residues. SEQ ID NO: 7192, includes amino acids 879 to 1005 of SEQ ID NO: 6042.

A preferred fragment of HR1 comprises amino acid residues 879 to 980 of SEQ ID NO: 6042. This preferred fragment is SEQ ID NO: 7193.

Another preferred fragment of HR1 comprises amino acid residues 901 to 1005 of SEQ ID NO: 6042. This preferred fragment is SEQ ID NO: 7194.

The SARS virus HR2 region comprises approximately amino acids 1144 to 1201 of SEQ ID NO: 6042, or fragments thereof capable of forming at least one alpha-helical turn. Preferably, said fragments comprise at least 7 (e.g., at least 14, 21, 28, 35, 42, 49 or 56) amino acid residues. SEQ ID NO: 7195 includes amino acids 1144 to 1201. A preferred fragment of HR2 comprises amino acids 1144 to 1195 of SEQ ID NO: 6042. This preferred fragment is SEQ ID NO: 7196.

Membrane Fusion peptides sequences within the spike protein are also believed to participate in fusion (and infection) of the virus with a host cell. Fusion peptides generally comprise about 16 to 26 amino acid residues which are conserved within viral families. These Membrane Fusion peptides are relatively hydrophobic and generally show an asymmetric distribution of hydrophobitity when modeled into an alpha helix. They are also generally rich in alanine and glycine.

At least three hydrophobic Membrane Fusion peptide regions have been identified within coronaviruses (PEP1, PEP2, and PEP3). See, Luo et al., "Roles in Cell-Cell Fusion of Two Conserved Hydrophobic Regions in the Murine Coronavirus Spike Protein", Virology (1998) 244:483-494. Figure 1 of this paper shows an alignment of Membrane Fusion peptide sequences of Mouse Hepatitis Viris, Bovine Corona Virus, Feline Infectious Peritonitis Virus, Transmissible Gastroenteritis Virus and Infectious Bronchitis Virus. See also, Bosch et al., "The Coronavirus Spike Protein is a Class I Virus Fusion Protein: Structural and Functional Characterization of the Fusion Core Complex" Journal of Virology (2003) 77(16):8801-8811.

PEP1 (SEQ ID NO: 7197), PEP2 (SEQ ID NO: 7198) and PEP3 (SEQ ID NO: 7199) sequences within the SARS spike protein have been identified.

The coronavirus spike proteins (and other similar surface viral proteins) are thought to undergo a conformational change upon receptor binding to the target cell membrane. One or more of the hydrophobic Membrane Fusion peptides are thought to become exposed and inserted into the target membrane as a result of this conformational change. The free energy released

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upon subsequent refolding of the spike protein to its most stable conformation is believed to play a role in the merger of the viral and cellular membranes.

One or more SARS HR sequences, preferably HR2, or a fragment thereof may be used to inhibit viral entry and membrane fusion with a target mammalian host cell. The invention provides a method of inhibiting viral infection comprising administering a composition comprising one or more SARS HR polypeptides or a fragment thereof. Preferably, the composition comprises a SARS HR2 sequence.

In another embodiment, the invention includes a composition comprising a SARS HR1 sequence, or a fragment thereof and a SARS HR2 sequence, or a fragment thereof. The HR1 and HR2 sequences may optionally be associated together in an oligomer. The composition may comprise the intermediate domain sequence between the HR1 and HR2 domains. The use of such an intermediate sequence may facilitate oligomerization or other structural interaction between the HR regions.

HR sequences for use in the invention may be produced recombinantly by methods known in the art. The SARS HR sequences may be modified to facilitate bacterial expression. In particular, the HR sequences may be modified to facilitate transport of the recombinant protein to the surface of the bacterial host cell. For example, leader sequences to a bacterial membrane protein may be added to the N terminus of the recombinant HR sequences. HR sequences for use in the invention may alternatively be produced by chemical synthesis by methods known in the art (see below).

As discussed in more detail later in the specification, Applicants have identified structural similarities between the SARS spike protein and the surface protein of Neisseria meningitidis, NadA (and other similar bacterial adhesion proteins). Another means of facilitating bacterial expression of HR sequences includes the addition of the stalk and/or anchor sequences of a NadA-like protein to the C-terminus of the recombinant HR sequences. Recombinant sequences containing the bacterial anchor sequence may preferably be prepared in outer membrane vesicles (the preparation of which is discussed in more detail later in the application). Recombinant sequences missing the bacterial anchor sequences may be secreted and isolated from the supernatant.

The invention includes a polypeptide sequence comprising a first sequence and a second sequence, wherein said first sequence comprises a leader sequence for a bacterial membrane protein and wherein said second sequence comprises a HR sequence of a coronavirus. Preferably, said first sequence comprises the leader sequence for a bacterial adhesin protein. More preferably, said bacterial adhesion protein is NadA. Preferably said second sequence comprises HR1, HR2 or both. In one embodiment, the second sequence comprises HR1, HR2 and the intermediate domain sequence present in the naturally occrding spike protein. For

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example, the second sequence may comprise a fragment of a coronavirus spike protein comprising the amino acids starting with the N-terminus of the HR1 region and ending with the C-terminus of the HR2 region.

The invention further includes a polypeptide sequence comprising a first, second, third and fourth sequence, wherein the first sequence comprises a leader sequence for a bacterial membrane protein; wherein said second sequence comprises a HR sequence of a coronavirus; wherein said third sequence comprises a stalk domain of a bacterial adhesion protein; and wherein said fourth sequence comprises an anchor domain of a bacterial adhesion protein. In one embodiment, the first sequence comprising the leader peptide sequence is removed. In another embodiment, the third sequence comprising the stalk domain is removed. In another embodiment, the fourth sequence comprising the anchor domain is removed.

The polypeptide sequences of the above described constructs may be linked together by means known in the art, including, for example, via glycine linkers.

Examples of constructs which may be used in such bacterial expression systems are shown in FIGURE 50. Polypeptide sequences of each of the constructs illustrated in FIGURE 50 are given as SEQ ID NOS: 7200 to 7206.

```
7200 Leader NadA (1-29) - HR1 (879-980) - 6Xgly - HR2 (1144-1195) - stalk+anchor NadA (88-405)
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Administration of one of more of these Membrane Fusion sequences may also interfere with the ability of a coronavirus to fuse to a host cell membrane. Accordingly, the invention includes an isolated polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 7197, SEQ ID NO: 7198 and SEQ ID NO: 7199. The invention further includes an isolated polypeptide comprising an amino acid sequence having sequence homology to an amino acid sequence selected from the group consisting of SEQ ID NO: 7197, SEQ ID NO: 7198 and SEQ ID NO: 7199.

Two or more of these SARS Membrane Fusion peptides can be combined together. The invention includes a composition comprising two SARS Membrane Fusion peptides wherein said peptides are selected from at least two of the amino acids selected from the group consisting of SEQ ID NO: 7197, SEQ ID NO: 7198and SEQ ID NO: 7199, or a sequence having sequence identity thereto.

Two or more of the SARS Membrane Fusion peptides may be linked together.

Accordingly, the invention includes a polypeptide comprising a first amino acid sequence and a

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⁷²⁰¹ Leader NadA (1-29) - HR1 (879-980) - 6Xgly - HR2 (1144-1196) - stalk NadA (88-351)

⁷²⁰² Leader NadA (1-29) - HR1 - HR2 (879-1196) - stalk+anchor NadA (88-405)

⁷²⁰³ Leader NadA (1-29) - HR1 - HR2 (879-1196)-stalk NadA (88-351) 7204 HR1 - HR2 (879-1196)-stalk NadA (88-351)-6xhis

⁷²⁰⁵ Leader NadA (1-29) - HR1 - HR2 (879-1196)-anchor NadA (351-405)

⁷²⁰⁶ Leader NadA (1-29) - HR1 - HR2 (879-1196)

second amino acid sequence, wherein said first and second amino acid sequences are selected from the group consisting of SEQ ID NO: 7197, SEQ ID NO: 7198 and SEQ ID NO: 7199, or a sequence having sequence identity thereto. Preferably, said first amino acid sequence and said second amino acid sequence are different SARS Membrane Fusion peptides, i.e., they are not the same.

The invention also includes a method of treating or preventing SARS virus infection comprising administering one or more of the SARS Membrane Fusion peptide compositions described above

As discussed above, the spike protein is capable of forming trimers. The invention further includes a polypeptide comprising SEQ ID NO: 6042 in trimeric form. The invention includes a composition comprising at least polypeptides wherein each polypeptide comprises at least the alpha-helical coiled region of a SARS virus spike protein. Preferably, the spike protein comprises SEQ ID NO: 6042 or a fragment thereof.

The invention further includes a composition comprising a SARS virus spike protein or a fragment thereof wherein said protein is associated with a transmembrane and wherein said fragment comprises the alpha-helical region of the SARS virus spike protein. Preferably, the composition comprises at least three SARS virus spike proteins or a fragment thereof, wherein the fragment comprises the alpha-helical region of the SARS virus spike protein.

The invention further includes an antibody which specifically binds to a trimeric form of SARS virus spike proteins. Preferably, the spike protein comprises SEQ ID NO: 6042 or a fragment thereof. The invention includes an antibody which specifically binds to a trimeric form of SARS virus spike proteins wherein said proteins are associated with a transmembrane.

The invention further includes an antibody which specifically binds to a monomeric form of SARS virus spike protein or a fragment thereof. Preferably, the antibody specifically binds to a monomeric form of SEQ ID NO: 6042 or a fragment thereof.

The invention further includes a small molecule which interferes with or disrupts the coiling of a SARS viral spike protein trimer.

The invention further includes an attenuated SARS virus for use as a vaccine wherein said attenuated virus contains a polynucleotide insertion, deletion or substitution which does not disrupt the trimeric conformation of the SARS virus spike protein. The invention further includes an attenuated SARS virus for use as a vaccine wherein said attenuated virus contains a polynucleotide insertion, deletion or substitution which does not disrupt the alpha-helical formation of the SARS virus spike protein.

The spike protein may be recombinantly produced. In one embodiment, the spike protein is expressed in virus like particles so that the protein is attached to a cell membrane. Such attachment may facilitate presentation of immunogenic epitopes of the spike protein. Preferably,

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the alpha-helical portion of the spike protein is associated with the cell membrane. Preferably, the spike proteins form a trimer within the transmembrane region of attachment.

Predicted N-glycosylation sites of SEQ ID NO: 6042 are identified below:

5	Position		Potential	Jury	NG1yc
,				agreement	result
	29 NYTQ	SEQ ID NO: 7207	0.7751	(9/9)	+++
	65 NVTG	SEQ ID NO: 7208	0.8090	(9/9)	+++
	109 NKSQ	SEQ ID NO: 7209	0.6081	(7/9)	+
	119 NSTN	SEQ ID NO: 7210	0.7039	(9/9)	++
10	158 NCTF	SEQ ID NO: 7211	0.5808	(7/9)	+
	227 NITN	SEQ ID NO: 7212	0.7518	(9/9)	+++
	269 NGTI	SEQ ID NO: 7213	0.6910	(9/9)	
	318 NITN	SEQ ID NO: 7214	0.6414	(9/9)	++
	330 NATK	SEQ ID NO: 7215	0.6063		++
15	357 NSTF	SEQ ID NO: 7216	0.5746	(8/9)	+
	589 NASS	SEQ ID NO: 7216		(8/9)	+
	602 NCTD		0.5778	(6/9)	+
	699 NFSI		0.6882	(9/9)	++
		SEQ ID NO: 7219	0.5357	(7/9)	+
20	783 NFSQ	SEQ ID NO: 7220	0.6348	(9/9)	++
20	1080 NGTS	SEQ ID NO: 7221	0.5806	(7/9)	+
	1116 NNTV	SEQ ID NO: 7222	0.5106	(5/9)	+
	1176 NESL	SEQ ID NO: 7223	0.6796		++

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment comprises one or more of the glycosylation sites identified above (SEQ ID NOS: 7207-7223). The invention further includes a polynucleotide encoding one or more of the fragments identified above. This glycosylation site can be covalently attached to a saccharide. Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment comprises one or more of the glycosylation sites identified above and wherein said polypeptide is glycosylated at one or more of the sites identified above.

Predicted O-glycosylation sites are identified below:

	Residu	e No.	Potential	Threshold Assig	mment
	Thr	698	0.8922	0.7696	т
_	Thr	706	0.9598	0.7870	Ť
5	Thr	922	0.9141	0.7338	Ť
	Ser	36	0.8906	0.7264	ŝ
	Ser	703	0.8412	0.7676	s

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment comprises one or more of the O-glycosylation sites identified above. The invention further includes a polynucleotide encoding one or more of the fragments identified above. The invention further includes a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment comprises one or more of the O-glycosylation sites identified above and further wherein the polypeptide is covalently bonded to a saccharide at one or more of the included glycosylation sites.

The invention further includes a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment comprises one or more of the N-glycosylation sites identified above and

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further wherein said fragment comprises one or more of the O-glycosylation sites identified above.

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polypucleotide encoding such a polypeptide.

Predicted phosphorylation sites of SEQ ID NO: 6042 are Ser-346, Tyr-195, and Tyr-723. Accordingly, the invention comprises a polypeptide comprising a fragment of SEQ ID NO: 6042 wherein said fragment comprises at least ten amino acid residues and wherein said fragment comprises one or more of the amino acids selected from the group consisting of Ser-346, Tyr-195, and Tyr-723. In one embodiment, one or more of the amino acids selected from the group consisting of Ser-346. Tyr-195, and Tyr-723 are phosphorylated.

Expression and functional characterization of the spike glycoprotein has been described by Xiao et al. (2003) Biochem Biophys Res Comm 312:1159-1164.

T-epitopes for SEQ ID NO: 6042 are identified in Table 16. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified as SEQ ID NOS: 8041-8280; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polypucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polypucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 8041-8280, or a polypucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide comprising SEQ ID NO: 6040. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 6040. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 6040.

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The invention includes a polynucleotide encoding SEQ ID NO: 6040 or a fragment thereof. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6040 or a fragment thereof. The invention includes a diagnostic kit comprising a polynucleotide sequence encoding SEQ ID NO: 6040 or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 6040 or a fragment thereof.

SEQ ID NO: 6040 demonstrates functional homology with a membrane protein of coronaviruses. Predicted transmembrane helices of SEQ ID NO: 6040 are identified below:

Predicted Transmembrane Helices

The sequence positions in brackets denominate the core region. Only scores above 500 are considered significant.

		~~		acar.		TIGITO	CO	-	Louisa
		:	Erom			to	score	CE	enter
15	27	(30)	48	(45)	1138		38
	137	(139)	153	(153)	486		146
	Outsi	de	e to	insid	le	helic	es :	3	found
			from			to	score	CE	enter
20						45)	819		38
	71	(73)	90	(90)	210		81
	136	(142)	156	(156)	272		149

Inside to outside belices .

The amino acid region with the highest predicted transmembrane helical region is from amino acid position 27 to 48 of SEQ ID NO: 6040. Such transmembrane regions are often difficult to express recombinantly. Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6040 wherein said fragment does not include the amino acid sequence between positions 27 to 48. The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6040 wherein said fragment does not include the amino acid sequence between positions 28 to 45. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6040 is predicted to be a hypothetical protein of the SARS virus. A prediction of the protein localization of SEQ ID NO: 6040 is set forth below. SEQ ID NO: 6040 is predicted to be located in one of the following locations: mitochondrial matrix space, microbody (peroxisome), nucleus, and mitochondrial inner membrane. SEQ ID NO: 6040 is predicted to be associated with an organelle inside an infected cell.

Accordingly, SEQ ID NO: 6040 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 6040 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6040 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6040 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6040 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEO

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ID NO: 6040 from associating with an organelle inside of an infected cell. The invention includes a fusion protein wherein said fusion protein comprises SEO ID NO: 6040.

```
PSORT --- Prediction of Protein Localization Sites
                                                    version 6.4(WWW)
5
       SEO ID NO: 6040
                                163 Residues
       Species classification: 4
       *** Reasoning Step: 1
10
       Preliminary Calculation of ALOM (threshold: 0.5)
            count: 0
       McG: Examining signal sequence (McGeoch)
            Length of UR:
                             9
            Peak Value of UR:
                                1.75
15
            Net Charge of CR: 1
            Discriminant Score:
                                    -2.56
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): 1.94
            Possible cleavage site: 53
20
       >>> Seems to have no N-terminal signal seg.
       Amino Acid Composition of Predicted Mature Form:
          calculated from 1
       ALOM new cnt: 0 ** thrshld changed to -2
       Cleavable signal was detected in ALOM?: OB
25
       ALOM: finding transmembrane regions (Klein et al.)
            count: 0 value: 1.32 threshold: -2.0 PERIPHERAL Likelihood = 1.32
            modified ALOM score: -1.16
       Gavel: Examining the boundary of mitochondrial targeting seq.
30
             motif at: 156
            HRSVTI
       Discrimination of mitochondrial target seq.:
            notclr ( 0.88)
       Rule: mitochondrial protein
       Rule: mitochondrial protein
35
       Rule: mitochondrial protein
       Rule: mitochondrial protein
       *** Reasoning Step: 2
40
             Count: 0
       Checking apolar signal for intramitochondrial sorting
         (Gavel position 156) from: 27 to: 44 Score: 5.0
       Mitochondrial matrix? Score: 0.36
45
       SKL motif (signal for peroxisomal protein):
            pos: 99(163), count: 1 SKL
            SKL score (peroxisome): 0.3
       Amino Acid Composition Tendency for Peroxisome: -4.28
       Peroxisomal proteins? Status: notclr
50
       Amino Acid Composition tendency for lysosomal proteins
            score: 0.02 Status: notclr
       Modified score for lysosome: 0.152
       Checking the amount of Basic Residues (nucleus)
       Checking the 4 residue pattern for Nuclear Targeting
55
            Found: pos: 132 (5) KRKR
       Checking the 7 residue pattern for Nuclear Targeting
       Checking the Robbins & Dingwall consensus (nucleus)
       Checking the RNA binding motif (nucleus or cytoplasm)
       nuc modified.
                       Score: 0.60
60
       Nuclear Signal
                        Status: notclr ( 0.30)
                                           -40-
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Checking CaaX motif..
Checking N-myristoylation..
Checking CaaX motif..
```

5 ---- Final Results ---mitochondrial matrix space --- Certainty= 0.480(Affirmative) < succ>
microbody (peroxisome) --- Certainty= 0.300(Affirmative) < succ>
nucleus --- Certainty= 0.300(Affirmative) < succ>
mitochondrial inner membrane --- Certainty= 0.188(Affirmative) < succ>

Predicted N-glycosylation sites of SEQ ID NO: 6040 are identified below.

	Pos	ition			Potential	Jury agreement	NGlyc result
15					0.7804 0.6123	(9/9) (7/9)	+++

Accordingly, the invention comprises a fragment of SEQ ID NO: 6040 wherein said fragment is at least ten amino acids and wherein said fragment comprises one or more of the asparagines from the amino acid positions of SEQ ID NO: 6040 selected from the group consisting of 2 and 106. The invention includes a fragment of SEQ ID NO: 6040 wherein said fragment comprises one or more amino acid sequences selected from the group consisting of SEQ ID NO: 7255 and SEQ ID NO: 7256. Preferably, the fragment comprises the amino acid sequence NKTG (SEQ ID NO: 7255).

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6040 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polypucleotide encoding such a polypeptide.

T-epitopes for SEQ ID NO: 6040 are identified in Table 14. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 7640-7800; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 7640-7800, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus.

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The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide comprising SEQ ID NO: 6041. SEQ ID NO: 6041 demonstrates functional homology with a portion of an ORF 1ab polyprotein. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 6041. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 6041. The invention includes a polynucleotide sequence encoding an amino acid sequence having sequence identity to SEQ ID NO: 6041. The invention includes a polynucleotide encoding a fragment of a polypeptide comprising SEQ ID NO: 6041.

The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6041 or a fragment therof. The invention includes a diagnostic kit comprising a polynucleotide encoding SEQ ID NO: 6041 or a fragment thereof. The invention includes an immunogenic composition comprising a polypeptide comprising SEQ ID NO: 6041 or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 6041 or a fragment thereof.

The polyproteins of coronaviruses are associated with enzymatic activity. Accordingly, SEQ ID NO: 6041 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprising SEQ ID NO: 6041 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6041 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6041 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6041 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEQ ID NO: 6041 from performing enzymative activity. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6041.

Predicted transmembrane or hydrophobic regions of SEQ ID NO: 6041 are identified below. Although the polyprotein of coronaviruses is proteolytically cleaved into numerous smaller proteins, hydrophobic domains in the polyprotein are known to mediate the membrane association of the replication complex and to be able to dramatically alter the architecture of host cell membranes. Accordingly, the hydrophobic domains of the polyprotein are targets for genetic mutation to develop attenuated SARS virus vaccines. The hydrophobic domains are also

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targets for small molecule inhibitors of the SARS virus. The hydrophobic domains may also be used to generate antibodies specific to those regions to treat or prevent SARS virus infection.

Possible transmembrane helices of SEQ ID NO: 6041

The sequence positions in brackets denominate the core region. Only scores above 500 are considered significant.

	Inside to outside hel	ices : 18 found
	from to	score center
	234 (234) 254 (250) 1046 241
10	256 (256) 272 (270) 252 263
	319 (319) 334 (334) 227 327
	503 (505) 522 (519) 405 512
	613 (615) 633 (629) 619 622
	677 (679) 703 (696) 467 689
15	849 (851) 869 (865)) 229 858
	1080 (1080)1097 (1094)) 306 1087
	1147 (1149)1163 (1163)) 354 1156
	1557 (1557)1581 (1577)) 817 1567
	1954 (1954)1971 (1971)	832 1964
20	2369 (2372)2395 (2387)) 300 2379
	2513 (2513)2532 (2529)) 690 2522
	D. L 1	
	Outside to inside hel	
25	from to	score center
43	239 (239) 254 (254)	
	239 (248) 272 (263)	
	311 (314) 334 (328)	
	499 (503) 522 (519)	
20	617 (617) 634 (631)	022
30	849 (853) 872 (872)	
	1147 (1147)1162 (1162)	
	L564 (1564)1581 (1579)	
	1951 (1951)1968 (1966)	
3.5	2513 (2522)2539 (2537)	711 2529
35		

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6041, wherein said fragment comprises an amino acid sequence including one or more of the hydrophobic transmembrane sequences identified above. The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6041 wherein said fragment comprises one or more of the following polypeptide sequences of SEQ ID NO: 6041: 234-254, 613-633, 1557-1581, 1954-1971, 2513-2532, 239-254, 1564-1581, 1951-1968, 2513-2539. Preferably, the fragment comprises one or more of the following polypeptide sequences of SEQ ID NO: 6041: 234-254 and 239-254. The invention also includes polynucleotides encoding each of the polypeptide fragments identified above.

The invention includes an attenuated SARS virus wherein said attenuated SARS virus contains an addition, deletion or substitution in the polynucleotides encoding for one of the hydrophobic domains identified above. The invention also includes a method for creating an attenuated SARS virus comprising mutating a SARS virus by adding, deleting or substituting the

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viral genome of the SARS virus to alter the coding of one or more of the hydrophobic domains of SEQ ID NO: 6041 identified above.

The invention includes an antibody which specifically identifies one or more of the hydrophobic regions of SEQ ID NO: 6041 identified above. The invention includes a small molecule which binds to, interferes with the hydrophobicity of or otherwise disrupts one or more of the hydrophobic regions of SEQ ID NO: 6041 identified above.

Predicted N-glycosylation sites of SEO ID NO: 6041 are identified below:

Position			Potential	Jury	NGlyc
				agreement	result
10	571 NLSH	(SEQ ID NO:	7257) 0.6598	(8/9)	+
	835 NTSR	(SEQ ID NO:	7258) 0.5762	(7/9)	+
	958 NVTD	(SEQ ID NO:	7259) 0.7494	(9/9)	++
	1113 NISD	(SEQ ID NO:	7260) 0.7259	(8/9)	+
	1205 NSTL	(SEQ ID NO:	7261) 0.6296	(9/9)	++
15	1460 NVTG	(SEQ ID NO:	7262) 0.6844	(9/9)	++
	1685 NHSV	(SEQ ID NO:	7263) 0.5181	(5/9)	+
	2029 NKTT	(SEQ ID NO:	7264) 0.5423	(5/9)	+

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEQ ID NO: 6041, wherein said fragment comprises one or more of the N-glycosylation sites identified above. The invention comprises a polypeptide comprising a fragment of SEQ ID NO: 6041 wherein said fragment comprises one or more of sequences SEQ ID NOS: 7257-7264. Preferably, the fragment comprises one or more of the sequences SEQ ID NOS: 7257, 7259, 7260, 7261 and 7262. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6041 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polypucleotide encoding such a polypeptide.

T-epitopes for SEQ ID NO: 6041 are identified in Table 15. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 7801-8040; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 7801-8040, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a

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CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus.

The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide sequence SEQ ID NO: 6043 or a fragment thereof. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 6043. The invention includes a polynucleotide sequence encoding the amino acid sequence of SEQ ID NO: 6043 or a fragment thereof.

Predicted transmembrane regions of SEQ ID NO: 6043 are set forth below.

	110	.	icica i	i ai isii	LCI.	iioranc	regions	or and m
15	Insid	lе	to o	utsid	le	helic	es :	4 found
		:	from			to	score	center
	41	(41)	56	(56)	1789	49
								89
	105	(105)	125	(125)	1250	115
20								
	Outsi	Ĺđ	e to	insid	le	helic	es :	3 found
		:	Erom			to	score	center
	41	(41)	59	(56)	2053	49
	76	(82)	98	(96)	1580	89
25	103	(105)	125	(123)	1257	115

The amino acid region with the highest predicted transmembrane helical region is from amino acid position 76 to 99 of SEQ ID NO: 6043. Such transmembrane regions are often difficult to express recombinantly. Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6043 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. Preferably, the fragment does not include the amino acids between positions 27 to 48. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6043 is predicted to be a hypothetical protein of the SARS virus. A prediction of the protein localization of SEQ ID NO: 6043 is set forth below. SEQ ID NO: 6043 is predicted to be located in one of the following locations: mitochondrial inner membrane, plasma membrane, Golgi body, and mitochondrial intermembrane space. SEQ ID NO: 6043 may be associated with an organelle inside an infected cell.

Accordingly, SEQ ID NO: 6043 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 6043 or a fragment thereof. The invention includes a polypucleotide encoding the polypeptide sequence of SEQ ID

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NO: 6043 or a fragment thereof. The invention includes a method of screening SEO ID NO: 6043 for an inhibitor. The invention includes the recombinant expression of SEO ID NO: 6043 in a host cell. The invention includes a small molecule which prevents the polypertide of SEO ID NO: 6043 from associating with an organelle inside of an infected cell. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6043.

```
PSORT --- Prediction of Protein Localization Sites for SEO ID NO: 6043
                                                   version 6.4(WWW)
       Species classification: 4
10
       *** Reasoning Step: 1
       Preliminary Calculation of ALOM (threshold: 0.5)
            count: 3
            Position of the most N-terminal TMS: 40 at i=2
15
       MTOP: membrane topology (Hartmann et al.)
            I(middle): 47
                          Charge diffirence(C-N): 3.5
       McG: Examining signal sequence (McGeoch)
            Length of UR:
                             12
            Peak Value of UR:
20
            Net Charge of CR: 0
            Discriminant Score:
                                    -4.67
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): 3.44
            Possible cleavage site: 15
25
       >>> Seems to have no N-terminal signal seg.
       Amino Acid Composition of Predicted Mature Form:
          calculated from 1
       ALOM new cnt: 2 ** thrshld changed to -2
       Cleavable signal was detected in ALOM?: OB
30
       ALOM: finding transmembrane regions (Klein et al.)
            count: 2 value: -6.90 threshold: -2.0
            INTEGRAL
                        Likelihood = -6.90
                                            Transmembrane
                                                             83 - 99 ( 78 - 101)
                        Likelihood = -5.04
                                                             40 - 56 (
                                                                         37 -
            INTEGRAL
                                             Transmembrane
            PERIPHERAL Likelihood = -0.32
35
            modified ALOM score:
                                   1.48
       >>> Likely a Type IIIb membrane protein (Nexo Ccyt)
       Gavel: Examining the boundary of mitochondrial targeting seq.
             motif at: 128
            MRCWLC
40
       Discrimination of mitochondrial target seg .:
            notclr ( 0.76)
       Rule: mitochondrial protein
       Rule: mitochondrial protein
       Rule: mitochondrial protein
45
       Rule: mitochondrial protein
       *** Reasoning Step: 2
       Type IIIa or IIIb is favored for ER memb. proteins
50
              Count: 0
       Checking apolar signal for intramitochondrial sorting
         (Gavel position 128) from: 39 to: 56 Score: 11.5
       >>> Seems to have an intramitochondrial signal
       Mitochondrial inner membrane? Score: 0.59
55
       Mitochondrial intermemb.space? Score: 0.22
       SKL motif (signal for peroxisomal protein):
            pos: 92(274), count: 1
                                     SHL
```

```
SKL score (peroxisome): 0.3
       Amino Acid Composition Tendency for Peroxisome:
       Peroxisomal proteins? Status: positive
       Amino Acid Composition tendency for lysosomal proteins
 5
            score: 1.16 Status: notclr
       Type III proteins may be localized at Golgi
       Checking the amount of Basic Residues (nucleus)
       Checking the 4 residue pattern for Nuclear Targeting
       Checking the 7 residue pattern for Nuclear Targeting
10
       Checking the Robbins & Dingwall consensus (nucleus)
       Checking the RNA binding motif (nucleus or cytoplasm)
       Nuclear Signal Status: negative ( 0.00)
       Check the Number of TMSs for typeIII (plasma memb.)
       Checking N-myristoylation..
15
       ---- Final Results ----
       mitochondrial inner membrane --- Certainty= 0.664(Affirmative) < succ>
       plasma membrane --- Certainty= 0.600(Affirmative) < succ>
       Golgi body --- Certainty= 0.400(Affirmative) < succ>
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       mitochondrial intermembrane space --- Certainty= 0.362(Affirmative) < succ>
```

Predicted N- and O- glycosylation sites of SEQ ID NO: 6043 are identified below.

							Tublitli
	Posit	ion			Potential	Jury	NGlyc
25	227 N	ATF (SEQ	ID NO:	7265)	0.6328	agreemen (7/9)	t result +
30	Resident Three Thr	due No. 28 32 34 170 267 268 269	Potent: 0.909 0.874 0.905 0.681 0.924 0.731	5 0 8 6 0 3	eshold Assi 0.6280 0.6595 0.6655 0.6600 0.5779 0.5708 0.5583	T T T T T	
35	Thr Ser Ser	270	0.802 0.693 0.645	3 .	0.5492 0.6091 0.5977	T S S	

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BNSDOCID: <WO____2004092360A2 | >

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEQ ID NO: 6043, wherein said fragment comprises the N-glycosylation sites or O-glycosylation sites identified above. The invention comprises a polypeptide comprising a fragment of SEQ ID NO: 6043 wherein said fragment comprises one or more of the N-glycosylation sites or O-glycosylation sites identified above. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6043 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polypucleotide encoding such a polypeptide.

T-epitopes for SEQ ID NO: 6043 are identified in Table 17. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 8281-8486; (b) an amino acid sequence having sequence identity to an amino acid

sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 8281-8486, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide comprising SEQ ID NO: 6044. The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6044 or a sequence having sequence identity to SEQ ID NO:206. The invention includes a polynucleotide encoding SEQ ID NO: 6044.

SEQ ID NO: 6044 is identified as a hypothetical protein. Predicted hydrophobic or transmembrane regions of SEQ ID NO: 6044 are identified below:

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6044 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. Preferably, the fragment does not include the amino acids between positions 1 to 19. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6044 is predicted to be a hypothetical protein of SARS virus. A prediction of the protein localization of SEQ ID NO: 6044 is set forth below. SEQ ID NO: 6044 is predicted to be located in one of the following locations: nucleus, mitochondrial matrix, lysosome (lumen),

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and microbody (peroxisome). SEQ ID NO: 6044 may be associated with an organelle inside an infected cell.

Accordingly, SEQ ID NO: 6044 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 6044 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6044 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6044 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6044 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEQ ID NO: 6044 from associating with an organelle inside of an infected cell. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6044.

10 PSORT --- Prediction of Protein Localization Sites for SEQ ID NO: 6044 version 6.4(WWW) 154 Residues Species classification: 4 1.5 *** Reasoning Step: 1 Preliminary Calculation of ALOM (threshold: 0.5) count: 0 20 McG: Examining signal sequence (McGeoch) Length of UR: Peak Value of UR: Net Charge of CR: 1 Discriminant Score: -7.97 25 GvH: Examining signal sequence (von Heijne) Signal Score (-3.5): -3.28 Possible cleavage site: 34 >>> Seems to have no N-terminal signal seq. Amino Acid Composition of Predicted Mature Form: 30 calculated from 1 ALOM new cnt: 0 ** thrshld changed to -2 Cleavable signal was detected in ALOM?: OB ALOM: finding transmembrane regions (Klein et al.) count: 0 value: 1.43 threshold: -2.0 35 PERIPHERAL Likelihood = 1.43 modified ALOM score: -1.19 Gavel: Examining the boundary of mitochondrial targeting seq. motif at: 151 FRKKOV 40 Discrimination of mitochondrial target seg .: notclr (-0.46) *** Reasoning Step: 2 45 Count: 0 Checking apolar signal for intramitochondrial sorting (Gavel position 151) from: 46 to: 50 Score: 5.0 Mitochondrial matrix? Score: 0.36 SKL motif (signal for peroxisomal protein): 50 pos: -1(154), count: 0 Amino Acid Composition Tendency for Peroxisome: Peroxisomal proteins? Status: notclr AAC score (peroxisome): 0.149 Amino Acid Composition tendency for lysosomal proteins

```
score: 0.81 Status: notclr
       Modified score for lysosome: 0.231
       Checking the amount of Basic Residues (nucleus)
       Checking the 4 residue pattern for Nuclear Targeting
5
            Found: pos: 134 (3) KHKK
       Checking the 7 residue pattern for Nuclear Targeting
       Checking the Robbins & Dingwall consensus (nucleus)
            Found: pos: 136 (3) KK VSTNLCTHSF RKKOV
       Final Robbins Score (nucleus): 0.60
10
       Checking the RNA binding motif (nucleus or cytoplasm)
       nuc modified.
                     Score: 0.90
       Nuclear Signal
                        Status: positive ( 0.70)
       Checking CaaX motif...
       Checking N-myristoylation..
15
       Checking CaaX motif...
       ---- Final Results ----
       nucleus --- Certainty= 0.880(Affirmative) < succ>
```

nucleus --- Certainty= 0.880(Affirmative) < succ> mitochondrial matrix space --- Certainty= 0.360(Affirmative) < succ> lysosome (lumen) --- Certainty= 0.231(Affirmative) < succ> microbody (peroxisome) --- Certainty= 0.149(Affirmative) < succ>

One predicted O-glycosylation site of SEQ ID NO: 6044 is identified at residue 4:

Residue No. Potential Threshold Assignment
Thr 4 0.6839 0.6484 T

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEQ ID NO: 6044, wherein said fragment comprises the O-glycosylation site identified above. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6044 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polypucleotide encoding such a polypeptide.

T-epitopes for SEQ ID NO: 6044 are identified in Table 18. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 8487-8665; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 8487-8665, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS

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virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above. wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide sequence comprising SEQ ID NO: 6045. The invention includes a polypeptide sequence comprising an amino acid sequence having sequence identity to SEQ ID NO: 6045. The invention includes a polypeptide sequence comprising a fragment of SEQ ID NO: 6045. The invention includes a polynucleotide sequence encoding any of these polypeptides.

SEO ID NO: 6045 demonstrates functional homology with the envelope or small membrane protein of coronaviruses. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6045 or a fragment thereof. The invention includes a diagnostic kit comprising a polynucleotide encoding SEQ ID NO: 6045 or a fragment thereof. The invention includes an immunogenic composition comprising SEQ ID NO: 6045 or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEO ID NO: 6045 or a fragment thereof.

Predicted transmembrane regions of SEO ID NO: 6045 are identified below:

```
Inside to outside helices :
                              1 found
      from
                  to
                        score center
  17 ( 19) 33 ( 33)
                         2881
Outside to inside helices :
                              1 found
      from
                        score center
                  t.o
  17 ( 17)
            34 ( 34)
                         2981
```

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6045 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. Preferably, the fragment does not include the amino acids between positions 17 to 34. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides. In one embodiment, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6045 wherein said fragment does not include amino acid 35 residues 1-34 of SEQ ID NO: 6045.

```
Predicted protein Localization Site of SEQ ID NO: 6045 is below.
PSORT --- Prediction of Protein Localization Sites for SEQ ID NO: 6045
                                               version 6.4(WWW)
Species classification: 4
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*** Reasoning Step: 1

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Preliminary Calculation of ALOM (threshold: 0.5)
            count: 2
            Position of the most N-terminal TMS: 17 at i=1
5
       MTOP: membrane topology (Hartmann et al.)
            I(middle): 24 Charge diffirence(C-N): 2.0
       McG: Examining signal sequence (McGeoch)
            Length of UR: 29
            Peak Value of UR: 3.40
10
            Net Charge of CR: -2
            Discriminant Score:
                                   13.07
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): 4.37
            Possible cleavage site: 32
15
       ... positive value of mtop ...
       >>> Seems to have an uncleavable N-term signal seg.
       Amino Acid Composition of Predicted Mature Form:
          calculated from 1
       ALOM new cnt: 1 ** thrshld changed to -2
20
       Cleavable signal was detected in ALOM?: OB
       ALOM: finding transmembrane regions (Klein et al.)
           count: 1 value: -15.12 threshold: -2.0
            INTEGRAL
                     Likelihood =-15.12
                                            Transmembrane 17 - 33 (
            PERIPHERAL Likelihood = 0.47
25
           modified ALOM score: 3.12
      >>> Seems to be a Type Ib (Nexo Ccyt) membrane protein
           The cytoplasmic tail is from 34 to 76 (44 Residues)
      Rule: vesicular pathway
      Rule: vesicular pathway
30
      Rule: vesicular pathway
       ( 6) or uncleavable?
      Gavel: Examining the boundary of mitochondrial targeting seq.
            motif at: 6
           Uncleavable? Ipos set to: 16
35
      Discrimination of mitochondrial target seq.:
           notclr ( 0.19)
      Rule: vesicular pathway
      Rule: vesicular pathway
      Rule: vesicular pathway
Ю
      *** Reasoning Step: 2
      > Relative position of the end of the tail: 44%
      Memb.protein with uncleavable signl is often at ER
15
      KDEL Count: 0
      Checking apolar signal for intramitochondrial sorting
        (Gavel position 16) from: 70 to: 99 Score: 21.5
      >>> Seems to have an intramitochondrial signal
      SKL motif (signal for peroxisomal protein):
ю
           pos: -1(76), count: 0
      Amino Acid Composition Tendency for Peroxisome: -4.11
      Peroxisomal proteins?
                             Status: negative
      Amino Acid Composition tendency for lysosomal proteins
           score: 0.68 Status: notclr
:5
      Checking the amount of Basic Residues (nucleus)
      Checking the 4 residue pattern for Nuclear Targeting
      Checking the 7 residue pattern for Nuclear Targeting
      Checking the Robbins & Dingwall consensus (nucleus)
      Checking the RNA binding motif (nucleus or cytoplasm)
n
      Nuclear Signal Status: negative ( 0.00)
      Check cytoplasmic tail for typeIb (plasma memb.)
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0

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Checking the NPXY motif.
Checking the YXRF motif.
Checking N-myristoylation.
```

---- Final Results ---plasma membrane --- Certainty= 0.730(Affirmative) < succ>

endoplasmic reticulum (membrane) --- Certainty= 0.640(Affirmative) < succ> endoplasmic reticulum (lumen) --- Certainty= 0.100(Affirmative) < succ> outside --- Certainty= 0.100(Affirmative) < succ> outside --- Certainty= 0.100(Affirmative) < succ>

Predicted N-glycosylation sites of SEQ ID NO: 6045 are identified at residues 48 and 66:

	Position	Potential	Jury	NGlyc			
			agreement	result			
1.5	48 NVSL	0.6514	(9/9)	++	(SEQ	ID NO:	7266)
15	66 NSSE	0.5880	(7/9)	+	(SEQ	ID NO:	7267)

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEQ ID NO: 6045, wherein said fragment comprises one or more of the N-glycosylation sites identified above. The invention comprises a polypeptide comprising a fragment of SEQ ID NO: 6045 wherein said fragment comprises one or more of the N-glycosylation sites identified above. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention includes a polypeptide comprising a fragment of SEQ ID NO: 6045 wherein said fragment does not include one or more of the glycosylation sites identified above. The invention also includes a polynucleotide encoding such a polypeptide.

T-epitopes for SEQ ID NO: 6045 are identified in Table 19. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 8666-8820; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 8666-8820, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide sequence comprising SEQ ID NO: 6046. The invention includes polypeptide sequences comprising an amino acid sequence having sequence identity to SEQ ID NO: 6046. The invention includes a polypeptide sequence comprising a fragment of SEQ ID NO: 6046. The invention includes a polypucleotide encoding one of these polypeptides.

SEQ ID NO: 6046 has functional homology with a matrix protein of a coronavirus. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6046 or a fragment thereof. The invention includes a diagnostic kit comprising a polynucleotide encoding SEQ ID NO: 6046 or a fragment thereof. The invention includes an immunogenic composition comprising SEQ ID NO: 6046 or a fragment thereof. The invention includes an antibody which recognizes a polypeptide comprising SEQ ID NO: 6046 or a fragment thereof.

Predicted transmembrane regions of SEQ ID NO: 6046 are identified below.

```
Inside to outside helices :
                                        3 found
              from
                           to
                                  score center
          21 (
                21)
                      38 (
                            36)
                                   2412
                                             29
20
          51 (
                53)
                      69 (
                            69)
                                   2645
                                             60
          74 (
                82)
                      96 (
                            96)
                                   2464
                                             89
       Outside to inside helices :
                                        3 found
              from
                                  score center
                           +0
25
          18 ( 21)
                      38 (
                            38)
                                   2363
                                             28
          52 (
                52)
                      67 (
                            67)
                                   2363
                                             60
                76)
                                             84
          76 (
                      95 (
                            92)
                                   2605
```

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6046 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. Preferably, the fragment does not include the amino acids between positions selected from the group consisting of 18 to 38, 52 to 67 and 76 to 95. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

Predicted protein localization of SEQ ID NO: 6046 is set forth below.

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```
McG: Examining signal sequence (McGeoch)
            Length of UR: 1
            Peak Value of UR:
            Net Charge of CR: -3
 5
            Discriminant Score:
                                     2.21
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): 4.29
            Possible cleavage site: 39
       ... positive value of mtop ...
10
       >>> Seems to have an uncleavable N-term signal seg.
       Amino Acid Composition of Predicted Mature Form:
          calculated from 1
       Cleavable signal was detected in ALOM?: OB
       ALOM: finding transmembrane regions (Klein et al.)
15
            count: 3 value: -7.64 threshold: 0.5
            INTEGRAL.
                       Likelihood = -7.64
                                             Transmembrane
                                                             21 - 37 ( 18 -
                                                                               391
            INTEGRAL.
                        Likelihood = -7.59
                                                             50 - 66 (
                                             Transmembrane
                                                                         43 -
                                                                               72)
            INTEGRAL
                        Likelihood = -5.04
                                             Transmembrane
                                                             79 - 95 (
                                                                         72 -
            PERIPHERAL Likelihood = 2.38
20
            modified ALOM score: 2.13
       >>> Likely a Type IIIb membrane protein (Nexo Ccyt)
       Rule: vesicular pathway
       Rule: vesicular pathway
       Rule: vesicular pathway
25
       (2) or uncleavable?
       Gavel: Examining the boundary of mitochondrial targeting seq.
             motif at: 2
            Uncleavable? Ipos set to: 12
       Discrimination of mitochondrial target seq.:
30
            negative (-4.16)
       Rule: vesicular pathway
       Rule: vesicular pathway
       Rule: vesicular pathway
35
       *** Reasoning Step: 2
       Type IIIa or IIIb is favored for ER memb. proteins
       Memb.protein with uncleavable signl is often at ER
              Count: 0
40
       Checking apolar signal for intramitochondrial sorting
       SKL motif (signal for peroxisomal protein):
            pos: -1(221), count: 0
       Amino Acid Composition Tendency for Peroxisome:
       Peroxisomal proteins? Status: notclr
45
       Amino Acid Composition tendency for lysosomal proteins
            score: 2.30 Status: positive
       Type III proteins may be localized at Golgi
       Checking the amount of Basic Residues (nucleus)
       Checking the 4 residue pattern for Nuclear Targeting
50
       Checking the 7 residue pattern for Nuclear Targeting
       Checking the Robbins & Dingwall consensus (nucleus)
       Checking the RNA binding motif (nucleus or cytoplasm)
       Nuclear Signal Status: negative ( 0.00)
       Check the Number of TMSs for typeIII (plasma memb.)
55
       Checking N-myristoylation..
       ---- Final Results ----
       endoplasmic reticulum (membrane) --- Certainty= 0.685(Affirmative) < succ>
       plasma membrane --- Certainty= 0.640(Affirmative) < succ>
60
       Golgi body --- Certainty= 0.460(Affirmative) < succ>
       endoplasmic reticulum (lumen) --- Certainty= 0.100(Affirmative) < succ>
```

One predicted N-glycosylation sites of SEQ ID NO: 6046 is identified at residue 4:

Prediction of N-glycosylation sites

```
Position Potential Jury NGlyc agreement result
4 NGTI 0.8430 (9/9) +++ (SEO ID NO: 7268)
```

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEQ ID NO: 6046, wherein said fragment comprises the N-glycosylation site identified above. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention further comprises a polypeptide comprising a fragment of amino acid sequence SEQ ID NO: 6046, wherein said fragment does not include the N-glycosylation site identified above. The invention includes a polynucleotide encoding such a fragment.

A variant of SEQ ID NO: 6046 that is included within the invention is SEQ ID NO: 9963. Compared to SEQ ID NO: 6046, this sequence has Val at residue 72 instead of Ala.

T-epitopes for SEQ ID NO: 6046 are identified in Table 20. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 8821-9018; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 8821-9018, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

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The invention includes a polypeptide sequence comprising SEQ ID NO: 6047 or a fragment thereof or an amino acid sequence having sequence identity thereto. Predicted transmembrane regions of SEQ ID NO: 6047 are identified below.

```
Inside to outside helices .
                               2 found
      from
                  to
                         score center
       10)
             29 ( 27)
                           729
                                   17
  21 (
       24)
             41 ( 41)
                           640
                                   34
Outside to inside helices :
                               2 found
      from
                  to
                         score center
             22 ( 19)
   4 (
        4)
                           874
                                   12
  22 (
       24) 41 ( 41)
                           499
                                   31
```

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6047 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. Preferably, the fragment does not include the amino acids between positions selected from the group consisting of 4 to 22 and 22 to 41. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6047 is predicted to be a hypothetical protein of SARS virus. A prediction of the protein localization of SEQ ID NO: 6047 is set forth below. SEQ ID NO: 6047 is predicted to be located in one of the following locations: plasma membrane, endoplasmic reticulum, Golgi body, and microbody (peroxisome). SEQ ID NO: 6047 may be associated with an organelle inside an infected cell or with viral entry to a host cell.

Accordingly, SEQ ID NO: 6047 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 6047 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6047 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6047 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6047 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEQ ID NO: 6047 from associating with an organelle inside of an infected cell or interacting with a host cell membrane. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6047. Predicted protein localization of SEQ ID NO: 6047 is set forth below.

```
35

PSORT --- Prediction of Protein Localization Sites

Species classification: 4

*** Reasoning Step: 1

40

Preliminary Calculation of ALOM (threshold: 0.5)

count: 1

Position of the most N-terminal TMS: 2 at i=1

MTOP: membrane topology (Hartmann et al.)

I (middle): 9 Charge diffirence(C-N): 0.5

MGG: Examining scimnal sequence (McGeoch)
```

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20

25

```
Length of UR:
            Peak Value of UR: 3.08
            Net Charge of CR: 0
            Discriminant Score:
                                     5.12
 5
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): -4.45
            Possible cleavage site: 34
       >>> Seems to have an uncleavable N-term signal seg.
       Amino Acid Composition of Predicted Mature Form:
10
          calculated from 1
       ALOM new cnt: 1 ** thrshld changed to -2
       Cleavable signal was detected in ALOM?: 0B
       ALOM: finding transmembrane regions (Klein et al.)
            count: 1 value: -2.44 threshold: -2.0
15
            INTEGRAL
                       Likelihood = -2.44
                                            Transmembrane 2 - 18 (
            PERIPHERAL Likelihood = 1.22
            modified ALOM score: 0.59
       >>> Seems to be a Type II (Ncyt Cexo) membrane protein
            The cytoplasmic tail is from 1 to 1 (1 Residues)
20
       Rule: vesicular pathway
       Rule: vesicular pathway
       Rule: vesicular pathway
       (5) or uncleavable?
       Gavel: Examining the boundary of mitochondrial targeting seg.
25
            motif at: 5
            Uncleavable? Ipos set to: 15
       Discrimination of mitochondrial target seq.:
           notclr ( 1.48)
       Rule: vesicular pathway
30
       Rule: vesicular pathway
       Rule: vesicular pathway
       *** Reasoning Step: 2
35
       Relative position of the cytoplasmic tail: 1%
           Larger value (>30%) is favared for ER memb. proteins
       Memb.protein with uncleavable signl is often at ER
       KDEL Count: 0
       Checking apolar signal for intramitochondrial sorting
        (Gavel position 15) from: 64 to: 93 Score: 30.0
       >>> Seems to have an intramitochondrial signal
       SKL motif (signal for peroxisomal protein):
           pos: -1(63), count: 0
      Amino Acid Composition Tendency for Peroxisome: 1.91
       Peroxisomal proteins? Status: notclr
           AAC score (peroxisome): 0.161
      Amino Acid Composition tendency for lysosomal proteins
           score: 0.04 Status: notclr
      Checking the consensus for Golgi
      Checking the consensus for Golgi
      Checking the cytoplasmic tail of type II. (Golgi)
      Checking the amount of Basic Residues (nucleus)
      Checking the 4 residue pattern for Nuclear Targeting
      Checking the 7 residue pattern for Nuclear Targeting
      Checking the Robbins & Dingwall consensus (nucleus)
      Checking the RNA binding motif (nucleus or cytoplasm)
      Nuclear Signal Status: negative ( 0.00)
      Check mitochondrial signal for typeII (plasma memb.)
      Type II is favored for plasma memb. proteins
      Checking the NPXY motif..
      Checking the YXRF motif..
```

10

15

60

:5

5

20

25

30

Checking N-myristoylation..

```
---- Final Results ----
Plasma membrane --- Certainty= 0.685(Affirmative) < succ>
endoplasmic reticulum (membrane) --- Certainty= 0.640(Affirmative) < succ>
Golgi body --- Certainty= 0.370(Affirmative) < succ>
microbody (peroxisome) --- Certainty= 0.161(Affirmative) < succ>
```

T-epitopes for SEQ ID NO: 6047 are identified in Table 21. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 9019-9131; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 9019-9131, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide comprising SEQ ID NO: 6048, a fragment thereof or an amino acid sequence having sequence identity thereto. Predicted transmembrane regions of SEQ ID NO: 6048 are identified below.

35	Inside to outside from 3 (3) 18 (100 (100) 117 (to score 18) 1857	center 10
	Outside to inside		
	from		center
	1 (1) 15 (8
40	100 (100) 117 (115) 3009	107

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6048 wherein said fragment does not include one or more of the hydrophobic amino acid

sequences identified above. Preferably, the fragment does not include the amino acids between positions selected from the group consisting of 1 to 15 and 100 to 117. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6048 is predicted to be a hypothetical protein of SARS virus. A prediction of the protein localization of SEO ID NO: 6048 is set forth below. SEQ ID NO: 6048 is predicted to be located in one of the following locations: plasma membrane, lysosome (membrane). microbody (peroxisome), and endoplasmic reticulum (membrane). SEQ ID NO: 6048 may be associated with an organelle inside an infected cell or may interact with a host cell plasma membrane during viral entry to the host cell.

Accordingly, SEQ ID NO: 6048 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 6048 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6048 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6048 for an inhibitor. The invention includes the recombinant expression of SEO ID NO: 6048 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEO ID NO: 6048 from associating with an organelle inside of an infected cell or prevents the polypeptide from associating with the cell membrane of a host cell. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6048. Predicted protein localization of SEO ID NO: 6048 is set forth below.

```
:0
      PSORT --- Prediction of Protein Localization Sites
                                                   version 6.4(www)
       Species classification: 4
       *** Reasoning Step: 1
:5
      Preliminary Calculation of ALOM (threshold: 0.5)
           Position of the most N-terminal TMS: 3 at i=2
      MTOP: membrane topology (Hartmann et al.)
           I(middle): 10
                          Charge diffirence(C-N): -2.5
      McG: Examining signal sequence (McGeoch)
           Length of UR:
                             13
           Peak Value of UR:
                                3.38
           Net Charge of CR: 1
5
           Discriminant Score:
                                   10.02
      GvH: Examining signal sequence (von Heijne)
           Signal Score (-3.5): 2.56
           Possible cleavage site: 15
      >>> Seems to have a cleavable N-term signal seq.
      Amino Acid Composition of Predicted Mature Form:
         calculated from 16
      ALOM new cnt: 2 ** thrshld changed to -2
      Cleavable signal was detected in ALOM?: 1B
      ALOM: finding transmembrane regions (Klein et al.)
           count: 1 value: -14.75 threshold: -2.0
           INTEGRAL
                       Likelihood =-14.75
                                           Transmembrane 101 - 117 ( 95 - 120)
           PERIPHERAL Likelihood = 6.63
           modified ALOM score:
                                  3.05
```

Э

3

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```
>>> Seems to be a Type Ia membrane protein
            The cytoplasmic tail is from 118 to 122 (5 Residues)
       Rule: vesicular pathway
       Rule: vesicular pathway
 5
       Rule: vesicular pathway
       (15) or uncleavable?
       Gavel: Examining the boundary of mitochondrial targeting seq.
             motif at: 15
            Uncleavable? Ipos set to: 25
10
       Discrimination of mitochondrial target seg.:
            notclr ( 0.73)
       Rule: vesicular pathway
       Rule: vesicular pathway
       Rule: vesicular pathway
15
       *** Reasoning Step: 2
              Count: 0
       Checking apolar signal for intramitochondrial sorting
20
         (Gavel position 25) from: 3 to: 12 Score: 8.5
       SKL motif (signal for peroxisomal protein):
            pos: -1(122), count: 0
       Amino Acid Composition Tendency for Peroxisome:
           AAC not from the N-term., score modified
25
       Peroxisomal proteins? Status: notclr
           AAC score (peroxisome): 0.115
       Amino Acid Composition tendency for lysosomal proteins
           score: -0.40 Status: negative
      GY motif in the tail of typeIa? (lysosomal)
30
      Checking the amount of Basic Residues (nucleus)
      Checking the 4 residue pattern for Nuclear Targeting
      Checking the 7 residue pattern for Nuclear Targeting
      Checking the Robbins & Dingwall consensus (nucleus)
      Checking the RNA binding motif (nucleus or cytoplasm)
35
      Nuclear Signal
                      Status: negative ( 0.00)
       Type Ia is favored for plasma memb. proteins
      Checking the NPXY motif..
      Checking the YXRF motif..
      Checking N-myristoylation..
Ю
      Checking GPI anchor ...
      >>> Seems to be GPI-anchored (0.85)
      ---- Final Results ----
      plasma membrane --- Certainty= 0.919(Affirmative) < succ>
15
      lysosome (membrane) --- Certainty= 0.200(Affirmative) < succ>
      microbody (peroxisome) --- Certainty= 0.115(Affirmative) < succ>
      endoplasmic reticulum (membrane) --- Certainty= 0.100(Affirmative) < succ>
```

T-epitopes for SEQ ID NO: 6048 are identified in Table 22. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 9132-9308; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further

comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 9132-9308, or a polypucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide comprising SEQ ID NO: 6049, a fragment thereof or an amino acid sequence having sequence identity thereto. Predicted transmembrane or hydrophobic regions of SEQ ID NO: 6049 are identified below.

```
Inside to outside helices: 1 found from to score center (2 13 (13) 30 (28) 3532 20

Outside to inside helices: 1 found from to score center (9 (11) 29 (26) 3395 19
```

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6049 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6049 is predicted to be a hypothetical protein of SARS virus. A prediction of the protein localization of SEQ ID NO: 6049 is set forth below. SEQ ID NO: 6049 is predicted to be located in one of the following locations: outside, microbody (peroxisome), endoplasmic reticulum (membrane) and endoplasmic reticulum (lumen). The highest ranking indicates that SEQ ID NO: 6049 is located on the outside of a cell. Accordingly, SEQ ID NO: 6049 may be a surface exposed protein.

Accordingly, SEQ ID NO: 6049 may be used in an immunogenic composition to raise an immune response against the SARS virus. It also may be used to generate antibodies specific to the SARS virus. Such antibodies may be used in a method of treatment or prevention of a SARS virus infection. Such antibodies may further be used in a diagnostic test to identify the presence or absence of SARS virus in a biological sample.

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The invention includes a polypeptide comprises SEQ ID NO: 6049 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6049 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6049 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6049 in a host cell. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6049. Predicted protein localization of SEQ ID NO: 6049 is set forth below.

```
PSORT --- Prediction of Protein Localization Sites
                                                    version 6.4(WWW)
       Species classification: 4
10
        *** Reasoning Step: 1
       Preliminary Calculation of ALOM (threshold: 0.5)
            count: 1
15
            Position of the most N-terminal TMS: 11 at i=1
       MTOP: membrane topology (Hartmann et al.)
            I(middle): 18
                           Charge diffirence(C-N): -2.0
       McG: Examining signal sequence (McGeoch)
            Length of UR:
                              24
20
            Peak Value of UR:
            Net Charge of CR: -2
            Discriminant Score:
                                    13.56
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): 0.52
25
            Possible cleavage site: 25
       >>> Seems to have a cleavable N-term signal seq.
       Amino Acid Composition of Predicted Mature Form:
          calculated from 26
       ALOM new cnt: 1 ** thrshld changed to -2
30
       Cleavable signal was detected in ALOM?: 1B
       ALOM: finding transmembrane regions (Klein et al.)
            count: 0 value: 14.80 threshold: -2.0
            PERIPHERAL Likelihood = 14.80
           modified ALOM score: -3.86
15
       Rule: vesicular pathway
       Rule: vesicular pathway
      Rule: vesicular pathway
       (2) or uncleavable?
      Gavel: Examining the boundary of mitochondrial targeting seg.
Ю
            motif at: 2
           Uncleavable? Ipos set to: 12
      Discrimination of mitochondrial target seq.:
           notclr ( 1.42)
      Rule: vesicular pathway
      Rule: vesicular pathway
      Rule: vesicular pathway
      *** Reasoning Step: 2
0
             Count: 0
      Number of Potential N-glycosylation Sites: 0
      Out: score 0.800
      Checking apolar signal for intramitochondrial sorting
        (Gavel position 12) from: 44 to: 73 Score: 30.0
5
      >>> Seems to have an intramitochondrial signal
      SKL motif (signal for peroxisomal protein):
           pos: -1(44), count: 0
```

```
Amino Acid Composition Tendency for Peroxisome:
                                                          9.47
            AAC not from the N-term., score modified
       Peroxisomal proteins?
                               Status: notclr
            AAC score (peroxisome): 0.320
5
       Amino Acid Composition tendency for lysosomal proteins
            score: -6.47 Status: negative
       Number of NX(S/T) motif: 0
       Checking the amount of Basic Residues (nucleus)
       Checking the 4 residue pattern for Nuclear Targeting
10
       Checking the 7 residue pattern for Nuclear Targeting
       Checking the Robbins & Dingwall consensus (nucleus)
       Checking the RNA binding motif (nucleus or cytoplasm)
       Nuclear Signal Status: negative ( 0.00)
       Checking CaaX motif..
15
       Checking N-myristoylation..
       Checking CaaX motif...
```

---- Final Results ---- outside --- Certainty= 0.820(Affirmative) < succ> microbody (peroxisome) --- Certainty= 0.320(Affirmative) < succ> endoplasmic reticulum (membrane) --- Certainty= 0.100(Affirmative) < succ> endoplasmic reticulum (lumen) --- Certainty= 0.100(Affirmative) < succ>

T-epitopes for SEQ ID NO: 6049 are identified in Table 23. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 9309-9437; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 9309-9437, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

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The invention includes a polypeptide comprising SEQ ID NO: 6050 or a fragment thereof or an amino acid sequence having sequence identity thereto. Predicted transmembrane or hydrophobic regions are identified below.

```
Inside to outside helices :
                                      1 found
5
             from
                         to
                                score center
         13 ( 15)
                    32 ( 30)
                                  558
       Outside to inside helices :
                                      1 found
             from
                         to
                                score center
10
         16 ( 16) 30 ( 30)
                                 364
                                          23
```

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6050 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

SEQ ID NO: 6050 is predicted to be a hypothetical protein of SARS virus. A prediction of the protein localization of SEQ ID NO: 6050 is set forth below. SEQ ID NO: 6050 is predicted to be located in one of the following locations: lysosome (lumen), mitochondrial matrix space, mitochondrial inner membrane, and mitochondrial intermembrane space. SEQ ID NO: 6050 may be associated with an organelle inside an infected cell during the viral replication cycle.

Accordingly, SEQ ID NO: 6050 is a target for screening of chemical inhibitors to the SARS virus. The invention includes a polypeptide comprises SEQ ID NO: 6050 or a fragment thereof. The invention includes a polynucleotide encoding the polypeptide sequence of SEQ ID NO: 6050 or a fragment thereof. The invention includes a method of screening SEQ ID NO: 6050 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6050 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEQ ID NO: 6050 from associating with an organelle inside of an infected cell or prevents the polypeptide from associating with the cell membrane of a host cell. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6050. Predicted protein localization of SEO ID NO: 6050 is set forth below.

```
PSORT --- Prediction of Protein Localization Sites
                                                   version 6.4(WWW)
                      84 Residues
      Species classification: 4
;5
      *** Reasoning Step: 1
      Preliminary Calculation of ALOM (threshold: 0.5)
            count: 0
0
      McG: Examining signal sequence (McGeoch)
           Length of UR:
           Peak Value of UR:
           Net Charge of CR: 2
           Discriminant Score:
                                    -5.73
      GvH: Examining signal sequence (von Heijne)
```

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Signal Score (-3.5): -0.12
            Possible cleavage site: 29
       >>> Seems to have no N-terminal signal seg.
       Amino Acid Composition of Predicted Mature Form:
 5
          calculated from 1
       ALOM new cnt: 0 ** thrshld changed to -2
       Cleavable signal was detected in ALOM?: OB
       ALOM: finding transmembrane regions (Klein et al.)
            count: 0 value:
                               8.43 threshold: -2.0
10
            PERIPHERAL Likelihood = 8.43
            modified ALOM score: -2.59
       Gavel: Examining the boundary of mitochondrial targeting seg.
             motif at: 61
            ARCWYL
15
       Discrimination of mitochondrial target seg.:
            positive (1.66)
       Rule: mitochondrial protein
       Rule: mitochondrial protein
       Rule: mitochondrial protein
20
       Rule: mitochondrial protein
       *** Reasoning Step: 2
              Count: 0
25
       Checking apolar signal for intramitochondrial sorting
         (Gavel position 61) from: 52 to: 58 Score: 6.0
       Mitochondrial matrix? Score: 0.38
      SKL motif (signal for peroxisomal protein):
            pos: -1(84), count: 0
30
       Amino Acid Composition Tendency for Peroxisome:
       Peroxisomal proteins?
                              Status: notclr
            AAC score (peroxisome): 0.263
       Amino Acid Composition tendency for lysosomal proteins
            score: 2.86 Status: positive
35
       Modified score for lysosome: 0.850
       Checking the amount of Basic Residues (nucleus)
       Checking the 4 residue pattern for Nuclear Targeting
       Checking the 7 residue pattern for Nuclear Targeting
       Checking the Robbins & Dingwall consensus (nucleus)
40
       Checking the RNA binding motif (nucleus or cytoplasm)
       Nuclear Signal
                        Status: negative ( 0.00)
       Checking Caax motif...
       Checking N-myristoylation..
       Checking CaaX motif..
45
       ---- Final Results ----
       lysosome (lumen) --- Certainty= 0.850(Affirmative) < succ>
       mitochondrial matrix space --- Certainty= 0.544(Affirmative) < succ>
       mitochondrial inner membrane --- Certainty= 0.266(Affirmative) < succ>
50
       mitochondrial intermembrane space --- Certainty= 0.266(Affirmative) < succ>
         One predicted N-glycosylation sites of SEQ ID NO: 6050 is identified at residue 43:
       Position Potential
                             Jury
                                     NGlyc
```

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEO ID NO: 6050 wherein said fragment comprises the N-glycosylation site

agreement result

(9/9)

0.6713

43 NVTI

(SEQ ID NO: 7269)

identified above. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention further comprises a polypeptide comprising a fragment of amino acid sequence SEQ ID NO: 6050 wherein said fragment does not include the N-glycosylation site identified above. The invention includes a polynucleotide encoding such a fragment.

T-epitopes for SEQ ID NO: 6050 are identified in Table 24. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 9438-9538; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 9438-9538, or a polynucleotide encoding such a polypeptide.

The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a polypeptide sequence comprising SEQ ID NO: 6051 or a fragment thereof or an amino acid sequence having sequence identity thereto. The invention includes a polypeptide sequence comprising SEQ ID NO: 6052 or a fragment thereof or an amino acid sequence having sequence identity thereto.

SEQ ID NO: 6051 and SEQ ID NO: 6052 demonstrate functional homology with a nucleocapsid protein of a coronavirus. The invention includes a diagnostic kit comprising a polypeptide comprising SEQ ID NO: 6051, SEQ ID NO: 6052 or a fragment thereof. The invention includes a diagnostic kit comprising a polynucleotide encoding SEQ ID NO: 6051, SEQ ID NO: 6052 or a fragment thereof. The invention includes an immunogenic composition comprising SEQ ID NO: 6051, SEQ ID NO: 6052 or a fragment thereof. The invention includes

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an antibody which recognizes a polypeptide comprising SEQ ID NO: 6051, SEQ ID NO: 6052 or a fragment thereof.

SEQ ID NO: 6051 is predicted to be phosphorylated at Ser-79; Thr-92; Ser-106; Thr-116; Thr-142: Ser-184: Ser-188: Ser-202: Ser-236: Thr-248: Ser-251: Ser-256: Thr-377.

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6051 wherein said fragment includes one or more of the amino acid residues of SEQ ID NO: 6051 selected from the group consisting of Ser-79; Thr-92; Ser-106; Thr-116; Thr-142; Ser-184; Ser-188; Ser-202; Ser-236; Thr-248; Ser-251; Ser-256; Thr-377. The invention further includes a polypeptide comprising a fragment of SEQ ID NO: 6051 wherein said fragment does not include one or more of the amino acid residues of SEQ ID NO: 6051 selected from the group consisting of Ser-79; Thr-92; Ser-106; Thr-116; Thr-142; Ser-184; Ser-188; Ser-202; Ser-236; Thr-248; Ser-251; Ser-256; Thr-377. Two further useful fragments of the N protein (e.g. for immunoassay) are SEQ ID NOS: 9783 & 9784, which are lysine-rich and can be used to distinguish the SARS virus from other coronaviruses.

Predicted transmembrane regions of SEQ ID NO: 6051 are identified below.

Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 6051 wherein said fragment does not include one or more of the hydrophobic amino acid sequences identified above. The invention also includes a polynucleotide sequence encoding any of the above-identified polypeptides.

Predicted protein localization of SEQ ID NO: 6051 is set forth below. SEQ ID NO: 6051 is predicted to be localized near the nucleus, lysosome (lumen), mitochondrial matrix space, and microbody (peroxisome). The highest ranking is for localization near the nucleus. Coronavirus nucleocapsid proteins are known to bind to viral RNA. Coronavirus nucleocapsid proteins are also thought to be important for cell mediated immunity. Accordingly, the invention includes a polynucleotide comprising SEQ ID NO: 6051. The invention further includes a viral vector or particle suitable for in vivo delivery of the polynucleotide sequence comprising a SARS virus nuceocapsid polynucleotide sequence or a fragment thereof. In one embodiment, the polynucleotide comprises SEQ ID NO: 6051 or a fragment thereof. The invention further includes a method for eliciting a cell mediated immune response comprising delivering a polynucleotide encoding a SARS virus nucleocapsid protein or a fragment thereof to a mammal. In one embodiment, the polynucleotide comprising SEQ ID NO: 6051 or a fragment thereof.

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The invention further includes a method of screening SEQ ID NO: 6051 for an inhibitor. The invention includes the recombinant expression of SEQ ID NO: 6051 in a host cell. The invention includes a small molecule which prevents the polypeptide of SEQ ID NO: 6051 from binding to SARS virus RNA during viral replication. The invention includes a fusion protein wherein said fusion protein comprises SEQ ID NO: 6051. Predicted protein localization of SEQ ID NO: 6051 is set forth below.

```
PSORT --- Prediction of Protein Localization Sites
                                                    version 6.4(WWW)
       Species classification: 4
10
       *** Reasoning Step: 1
       Preliminary Calculation of ALOM (threshold: 0.5)
             count: 0
15
       McG: Examining signal sequence (McGeoch)
            Length of UR: 3
             Peak Value of UR:
            Net Charge of CR: 0
            Discriminant Score:
                                   -15.98
20
       GvH: Examining signal sequence (von Heijne)
            Signal Score (-3.5): -6.36
            Possible cleavage site: 58
       >>> Seems to have no N-terminal signal seq.
       Amino Acid Composition of Predicted Mature Form:
25 .
          calculated from 1
       ALOM new cnt: 0 ** thrshld changed to -2
       Cleavable signal was detected in ALOM?: 0B
       ALOM: finding transmembrane regions (Klein et al.)
            count: 0 value:
                               5.04 threshold: -2.0
30
            PERIPHERAL Likelihood = 5.04
            modified ALOM score: -1.91
       Gavel: Examining the boundary of mitochondrial targeting seq.
             motif at: 17
            PRITEG
35
       Discrimination of mitochondrial target seq.:
            negative (-3.97)
       *** Reasoning Step: 2
10
              Count: 0
       Checking apolar signal for intramitochondrial sorting
       Mitochondrial matrix? Score: 0.10
       SKL motif (signal for peroxisomal protein):
            pos: -1(399), count: 0
15
      Amino Acid Composition Tendency for Peroxisome:
       Peroxisomal proteins?
                              Status: notclr
            AAC score (peroxisome): 0.072
      Amino Acid Composition tendency for lysosomal proteins
           score: 0.96 Status: notclr
60
      Modified score for lysosome: 0.246
      Checking the amount of Basic Residues (nucleus)
      Checking the 4 residue pattern for Nuclear Targeting
           Found: pos: 256 (4) KKPR
           Found: pos: 372 (5) KKKK
i5
      Checking the 7 residue pattern for Nuclear Targeting
      Checking the Robbins & Dingwall consensus (nucleus)
           Found: pos: 372 (3) KK KKTDEAQPLP QRQKK
```

```
Found: pos: 373 (3) KK KTDEAOPLPO ROKKO
       Final Robbins Score (nucleus): 0.80
       Checking the RNA binding motif (nucleus or cytoplasm)
       nuc modified.
                       Score: 0.90
 5
       Nuclear Signal
                         Status: positive ( 0.90)
       Checking CaaX motif..
       Checking N-myristoylation..
       Checking CaaX motif...
10
       ---- Final Results ----
       nucleus --- Certainty= 0.980(Affirmative) < succ>
       lysosome (lumen) --- Certainty= 0.246(Affirmative) < succ>
       mitochondrial matrix space --- Certainty= 0.100(Affirmative) < succ>
       microbody (peroxisome) --- Certainty= 0.072(Affirmative) < succ>
15
    Predicted N-glycosylation sites of SEO ID NO: 6051 are identified below.
       Position Potential
                              Jury
                                      NGlvc
```

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agreement result
        48 NNTA
                   0.6879
                              (9/9)
                                       ++
                                               (SEQ ID NO: 7270)
20
       270 NVTO
                   0.7684
                              (9/9)
                                               (SEQ ID NO: 7271)
       Residue No.
                    Potential Threshold Assignment
        Thr 166
                      0.8547
                                0.6439
                                          т
        Thr
             367
                      0.5575
                                0.5403
                                          Τį
25
        Thr 394
                      0.8217
                                0.5821
                                          т
```

Accordingly, the invention comprises a polypeptide comprising a fragment of the amino acid sequence of SEQ ID NO: 6051 wherein said fragment comprises one or more of the N-glycosylation sites identified above. The invention further comprises a polynucleotide encoding one or more of the polypeptides identified above.

The invention further comprises a polypeptide comprising a fragment of amino acid sequence SEQ ID NO: 6051 wherein said fragment does not include one or more of the N-glycosylation sites identified above. The invention includes a polynucleotide encoding such a fragment.

T-epitopes for SEQ ID NO: 6052 are identified in Table 25. The invention includes a polypeptide for use as an antigen, wherein the polypeptide comprises: (a) an amino acid sequence selected from the group consisting of the T-epitope sequences identified in SEQ ID NOS: 9539-9752; (b) an amino acid sequence having sequence identity to an amino acid sequence of (a). The invention further comprising a polynucleotide sequence encoding the polypeptides of (a) or (b). The invention further comprising a method of expression or delivery of such polynucleotides through viral vectors and/or viral particles. The invention further comprises a polypeptide comprising two or more of the T-epitope sequences identified in SEQ ID NOS: 9539-9752, or a polynucleotide encoding such a polypeptide.

A variant of SEQ ID NO: 6052 that is included within the invention is SEQ ID NO: 9964.

Compared to SEQ ID NO: 6052, this sequence has Ile at residue 54 instead of Thr.

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The use as an antigen is preferably a use: (1) as a T-cell antigen; (2) for generating a complex between a class I MHC protein (e.g. a class I HLA) and a fragment of said antigen; (3) as an antigen for raising a cell-mediated immune response; and/or (4) as an antigen for raising a CTL response. The use preferably protects or treats disease and/or infection caused by a SARS virus. The invention provides the use of a polypeptide in the manufacture of a medicament for immunising a mammal (typically a human) against SARS viral infection wherein the polypeptide is as defined above.

The invention provides a method of raising an immune response in a mammal (typically a human), comprising the step of administering to the mammal a polypeptide as defined above, wherein said immune response is a cell-mediated immune response and, preferably, a CTL response. The immune response is preferably protective or therapeutic.

The invention includes a composition comprising a SARS virus nucleocapsid protein or a fragment thereof and further comprising a SARS virus membrane protein or a fragment thereof. The composition may further comprising one or more adjuvants discussed below.

The invention further includes a composition comprising a polypeptide comprising SEQ ID NO: 6051 or a fragment thereof or a sequence having sequence identity thereto and further comprising a polypeptide comprising SEQ ID NO: 6040, or a fragment thereof or a sequence having sequence identity thereto. Such composition may be used, for instance, in a vaccine. Such composition may further comprise one or more adjuvants discussed below.

The invention includes a composition comprising a SARS virus nucleocapsid protein or a fragment thereof and a SARS virus spike protein or a fragment thereof. In one embodiment the nucleocapsid protein comprises a polypeptide sequence comprising SEQ ID NO: 6051 or a fragment thereof or a sequence having sequence identity thereto. In one embodiment, the spike protein comprises a polynucleotide comprising SEQ ID NO: 6042 or a fragment thereof or a sequence having sequence identity thereto. The composition may further comprise one or more of the adjuvants discussed below.

The invention further includes a composition comprising antibodies specific to a SARS virus nucleocapsid protein and comprising antibodies specific to a SARS virus spike protein. In one embodiment the antibody is specific to a nucleocapsid protein comprises a polypeptide sequence comprising SEQ ID NO: 6051 or a fragment thereof or a sequence having sequence identity thereto. In one embodiment, the antibody is is specific to a spike protein comprises a polynucleotide comprising SEQ ID NO: 6042 or a fragment thereof or a sequence having sequence identity thereto.

The invention further includes polynucleotide sequences, and fragments thereof, of a SARS virus which are conserved among coronaviruses, and polypeptides encoded thereby. Such conserved sequences can be identified in the alignments shown in FIGURE 7. Such conserved

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sequences may be used in the vaccines of the invention or in the diagnostic reagents, kits and methods of the invention.

The invention further includes polynucleotide sequences, and fragments thereof, of a SARS virus which are specific to SARS virus and not shared with coronaviruses. Such SARS specific sequences are also identified as SEQ ID NOS: 6040, 6043, 6044, 6047, 6048, 6049 and 6050. Such SARS specific sequences may be used in the vaccines of the invention or in the diagnostic reagents, kits and methods of the invention.

The invention also includes polynucleotide sequences which can be used as probes or primers for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in SEQ ID NOS: 6076-6265 (Table 5). The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in SEQ ID NOS: 6076-6265.

The invention also includes polynucleotide sequences which can be used as probes or primers for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in SEQ ID NOS: 6266-6343 (Table 6). The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in SEQ ID NOS: 6266-6343.

The invention also includes polynucleotide sequences which can be used as probes or primers for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in SEQ ID NOS: 6344-6392 (Table 7). The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in SEQ ID NOS: 6344-6392...

The invention also includes polynucleotide sequences which can be used as probes or primers for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in SEQ ID NOS: 6393-6559 (Tables 8 & 9). The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in SEQ ID NOS: 6393-6559.

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The invention also includes polynucleotide sequences which can be used as probes or primers for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer and probe sequences identified in SEQ ID NOS: 6560-6568. The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in SEQ ID NOS: 6560-6568.

The invention includes a polypeptide sequence comprising any one of even-numbered SEQ ID NOS: 7272-7290, or a fragment thereof, or a sequence having sequence identity thereto. The invention further includes a polynucleotide sequence encoding any one of even-numbered SEQ ID NOS: 7272-7290, or a fragment thereof, or a sequence having sequence identity thereto. Examples of such polynucleotide sequences are odd-numbered SEQ ID NOS: 7273-7291.

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BNSDOCID: <WO_____2004092360A2_I_>

The invention includes a polynucleotide sequence comprising an intergenic sequence which is common to each open reading frame of the SARS virus. The SARS virus is thought to use this sequence to signal translation of the open reading frame. The intergenic sequence comprises a 10mer SEQ ID NO: 7292, or optionally a hexamer SEQ ID NO: 7293. When the virus transcribes its positive (+) RNA strand to (-) RNA strand, the virus replicating structure uses the (-) strand template to transcribe nucleotides at the 5' end prior to the first intergenic sequence, followed by the intergenic sequence, followed by the selected open reading frame. The virus then creates multiple mRNAs comprising the 5' end, the intergenic sequence and coding sequence. For more details on Nidovriales replication (including Coronavirus) see e.g., Ziebuhr et al., "Virus-encoded proteinases and proteolytic processing in the Nidovirales", Journal of General Virology 81:853-879 (2000), incorporated herein by reference in its entirety.

The invention comprising a polynucleotide sequence comprising SEQ ID NO: 7292 or the complement thereof. The invention comprising a polynucleotide sequence comprising SEQ ID NO: 7293 or the complement thereof. The invention further comprises a polynucleotide sequence comprising nucleotides from the 5' end of the SARS viral genome, or its reverse complement, and further comprising an intergenic sequence or its reverse complement. The polynucleotide may further comprise one or more of the SARS virus open reading frames. Examples of polynucleotide sequences comprising nucleotides from the 5' end of the SARS virus genome followed by the intergenic sequence are SEO ID NOS: 7294-7301.

The invention includes a polynucleotide sequence comprising a sequence selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301, or a fragment thereof, or a sequence having sequence identity

thereto. In one embodiment, the polynucleotide does not consist entirely of a known SARS virus sequence.

The SARS virus intergenic sequence can be used to create a RNAi molecule. Such a SARS virus specific RNAi molecule can be used to treat SARS virus infection. The invention includes a RNAi molecule comprising a double stranded RNA molecule wherein one RNA strand comprises a sequence selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301, or a fragment thereof. Preferably, said RNA strand comprises a sequence selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293. Preferably, the other RNA strand comprises the reverse complement of the first strand or a polynucleotide sequence which hybridizes to the first strand.

The invention includes the use of RNAi in a method of treatment for SARS virus infection comprising administering to a mammal an effective amount of the si RNA molecule. Preferably, the RNAi molecule comprises the molecule described above. Further discussion of the RNAi applications of the intergenic sequence is included in section IV of the specification below.

The invention also includes the use of a SARS virus antisense nucleotide sequence, preferably antisense directed to the SARS virus intergenic sequence. Such an antisense sequence may be used in the treatment of a subject infected with the SARS virus. The antisense of the SARS virus intergenic sequence can be designed to bind to the SARS viral polynucleotides to block access of the viral replication machinery to the intergenic sequence. Such an antisense sequence may also be used to identify the presence or absence of a SARS virus in a biological sample. The antisence can itself be labeled or the antisense associated with viral polynucleotides can be detected by means known in the art.

Antisense nucleic acids are designed to specifically bind to RNA, resulting in the formation of RNA-DNA or RNA-RNA hybrids, with an arrest of DNA replication, reverse transcription or messenger RNA translation. Antisense polynucleotides based on a selected sequence can interfere with expression of the corresponding gene. Antisense polynucleotides will bind and/or interfere with the translation of the corresponding mRNA.

The invention also includes the use of the intergenic region with a ribozyme.

Trans-cleaving catalytic RNAs (ribozymes) are RNA molecules possessing endoribonuclease activity. Ribozymes are specifically designed for a particular target, and the target message must contain a specific nucleotide sequence. They are engineered to cleave any RNA species site-specifically in the background of cellular RNA. The cleavage event renders the mRNA unstable and prevents protein expression. Importantly, ribozymes can be used to inhibit

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expression of a gene of unknown function for the purpose of determining its function in an in vitro or in vivo context, by detecting the phenotypic effect.

One commonly used ribozyme motif is the hammerhead, for which the substrate sequence requirements are minimal. Design of the hammerhead ribozyme is disclosed in Usman et al., Current Opin. Struct. Biol. (1996) 6:527-533. Usman also discusses the therapeutic uses of ribozymes. Ribozymes can also be prepared and used as described in Long et al., FASEB J. (1993) 7:25; Symons, Ann. Rev. Biochem. (1992) 61:641; Perrotta et al., Biochem. (1992) 31:16-17; Ojwang et al., Proc. Natl. Acad. Sci. (USA) (1992) 89:10802-10806; and US Patent 5,254,678. Ribozyme cleavage of HIV-I RNA is described in US Patent 5,144,019; methods of cleaving RNA using ribozymes is described in US Patent 5,116,742; and methods for increasing the specificity of ribozymes are described in US Patent 5,225,337 and Koizumi et al., Nucleic Acid Res. (1989) 17:7059-7071. Preparation and use of ribozyme fragments in a hammerhead structure are also described by Koizumi et al., Nucleic Acids Res. (1989) 17:7059-7071.

Preparation and use of ribozyme fragments in a hairpin structure are described by Chowrira & Burke, Nucleic Acids Res. (1992) 20:2835. Ribozymes can also be made by rolling transcription as described in Daubendiek & Kool. Nat. Biotechnol. (1997) 15(3):273-277.

The hybridizing region of the ribozyme may be modified or may be prepared as a branched structure as described in Horn & Urdea, Nucleic Acids Res. (1989) 17:6959-67. The basic structure of the ribozymes may also be chemically altered in ways familiar to those skilled in the art, and chemically synthesized ribozymes can be administered as synthetic oligonucleotide derivatives modified by monomeric units. In a therapeutic context, liposome mediated delivery of ribozymes improves cellular uptake, as described in Birikh et al., Eur. J. Biochem. (1997) 245:1-16.

Therapeutic and functional genomic applications of ribozymes proceed beginning with knowledge of a portion of the coding sequence of the gene to be inhibited. In the present-invention, the target sequence preferably comprises the intergeneic sequence of the SARS virus. Preferably, the sequence is selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293. A target cleavage site is selected in the target sequence, and a ribozyme is constructed based on the 5' and 3' nucleotide sequences that flank the cleavage site. Preferably, the 5' nucleotide sequence includes the 5' untranslated region of the SARS virus. The ribozyme may then further be constructed from one or more of the polynucleotide sequences selected from the group consisting of SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7301.

Antisense treatment of HIV infection is described in the following references, each of which is incorporated herein by reference in their entirety. (antisense RNA complementary to the mRNA of gag, tat, rev, env) (Sezakiel et al., 1991, J. Virol. 65:468-472; Chatterjee et al.,

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1992, Science 258:1485-1488; Rhodes et al., 1990, J. Gen. Virol. 71:1965. Rhodes et al., 1991, AIDS 5:145-151; Sezakiel et al., 1992, J. Virol. 66:5576-5581; Joshi et al., 1991, J. Virol. 65:5524-5530).

The invention includes the use of decoy RNA to disrupt the SARS virus replication and life cycle. Methods of making and using such decoy RNA for treatment of a viral infection are known in the art. The invention includes delivery of genes encoding, for example, the SARS virus intergenic sequence, to infected cells. Preferably, the sequence comprises one or more of the sequences selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301. Preferably, the sequence comprises one or more of the sequences selected from the group consisting of SEQ ID NO: 7292 and SEO ID NO: 7293. Preferably, the sequence comprises SEO ID NO: 7293.

In the present invention, delivery of intergenic sequence which is not linked to the SARS virus open reading frames disrupts the translation process of the viral RNA and decreases the production of vial proteins. Similar methods of treatment for HIV viral infection have been described. The following references discuss the use of decoy RNA of HIV TAR or RRE for treatment of HIV infection. Each of these references is incorporated herein by reference in their entirety. (Sullenger et al., 1990, Cell 63:601-608; Sullenger et al., 1991, J. Virol. 65:6811-6816; Lisziewicz et al., 1993, New Biol. 3:82-89; Lee et al., 1994, J. Virol. 68:8254-8264), ribozymes (Sarver et al., 1990, Science 247:1222-1225; Wecrasinghe et al., 1991, J. Virol. 65:5531-5534; Dropulic et al., 1992, J. Virol. 66:1432-1441; Ojwang et al., 1992, Proc. Natl. Acad. Sci. USA. 89:10802-10806; Yu et al., 1993, Proc. Natl. Acad. Sci. USA. 90:6340-6344; Yu et al., 1995, Proc. Natl. Acad. Sci. USA. 92:699-703; Yamada et al., 1994, Gene Therapy 1:38-45).

The invention includes the use of the SARS virus intergenic sequence in diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. Such diagnostic reagents, kits, and methods are further discussed in Section II of the specification.

The invention includes a pair of primers for amplifying a SARS polynucleotide sequence comprising (i) a first primer comprising a sequence which is substantially identical to a portion of a sequence selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301 and (ii) a second primer comprising a sequence which is substantially complementary to a portion of a sequence selected from the group consisting of the sequence SEQ ID NO: 1 and the sequence SEQ ID NO: 2, such that the primer pair (i) and (ii) defines a template sequence within a sequence from the group consisting of the sequence SEQ ID NO: 1 and the sequence SEQ ID NO: 2. Preferably, the (i)

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first primer comprises a sequence which is substantially identical to a portion of a sequence selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293. Preferably, the (i) first primer comprises a sequence which is substantially identical to a portion of the sequence of SEQ ID NO: 7293. The amplicon defined by said first and second primers is preferably between 50 and 250 nucleotides in length. The primers may optionally be labeled to facilitate their detection. Methods and compositions for use in labeling primers are discussed further in the application in Section III.

The invention further includes a pair of primers for amplifying a SARS polynucleotide sequence comprising (i) a first primer comprising a sequence which is substantially identical to a portion of the complement of a portion of a sequence selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301 and (ii) a second primer comprising a sequence which is substantially complementary to a portion of the complement of a sequence selected from the group consisting of the sequence SEQ ID NO: 1 and the sequence SEQ ID NO: 2, such that the primer pair defines a template sequence within a sequence selected from the group consisting of the sequence SEQ ID NO: 1 and the sequence SEQ ID NO: 2. The amplicon defined by said first and second primers is preferably between 50 and 250 nucleotides in length. The primers may optionally be labeled to facilitate their detection. Methods and compositions for use in labeling primers are discussed further in the application in Section III.

The invention includes a kit comprising (i) a first primer comprising a sequence which is substantially identical to a portion of a sequence selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301 and (ii) a second primer comprising a sequence which is substantially complementary to a portion of a sequence selected from the group consisting of the sequence SEQ ID NO: 1 and the sequence SEQ ID NO: 2, such that the primer pair (i) and (ii) defines a template sequence within a sequence from the group consisting of SEQ ID NO: 1 and SEQ ID NO: 2. Preferably, the (i) first primer comprises a sequence which is substantially identical to a portion of a sequence selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293. Preferably, the (i) first primer comprises a sequence which is substantially identical to a portion of the sequence of SEQ ID NO: 7293. The primers may optionally be labeled to facilitate their detection. Methods and compositions for use in labeling primers are discussed further in the application in Section III.

Other preferred kits comprise (i) a first primer comprising a sequence which is substantially identical to a portion of the complement of a portion of a sequence selected from

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the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301 and (ii) a second primer comprising a sequence which is substantially complementary to a portion of the complement of a sequence selected from the group consisting of SEQ ID NO: 1 and SEQ ID NO: 2, such that the primer pair defines a template sequence within a sequence selected from the group consisting of SEQ ID NO: 1 and SEO ID NO: 2.

The invention further includes an attenuated SARS virus for use as a vaccine wherein the intergenic region has been mutated to reduce expression of the viral structural or nonstructural proteins. The attenuated SARS virus may comprises one or more additions, deletions or insertion in one or more of the intergenic regions of the viral genome. Preferably, the attenuated SARS virus comprises an addition, deletion or insertion in one or more occurrences of the sequence selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293.

Preferably, the addition, deletion or insertion occurs in one or more occurrences of SEQ ID NO: 7293.

The invention further comprises a small molecule which inhibits binding or association of the SARS viral replication machinery, such as a ribonucleoprotein, with the intergenic region of the viral genome. Preferably, the small molecule inhibits binding or association of the SARS viral machinery with a sequence selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293. Preferably, the small molecule inhibits binding or association of the SARS viral machinery with SEQ ID NO: 7293. The invention further includes a method of screening for a small molecule for treatment of SARS viral infection comprising using an assay to identify a small molecule which interferes with the association of the SARS viral replication machinery with the intergenic region of the SARS viral genome.

The invention further provides a novel SARS polynucleotide sequence SEQ ID NO: 9968. All six reading frames of this 690mer sequence are shown in Figure 113. The constituent amino acid sequences from Figure 113, having at least 4 amino acids, are listed as SEQ ID NOS: 9969 to 10032.

Accordingly the invention includes a polynucleotide sequence comprising SEQ ID NO: 9968. It also provides polynucleotide sequences having sequence identity to SEQ ID NO: 9968. The degree of sequence identity is preferably greater than 50% (e.g., 60%, 70%, 80%, 85%, 88%, 90%, 92%, 95%, 99% or more).

The invention includes an amino acid sequence encoded by the polynucleotide sequence of SEQ ID NO: 9968, including the amino acid sequences selected from the group consisting of SEQ ID NO^S: 9969 to 10032. Preferably, the amino acid sequence comprises SEQ ID NO: 9997 or comprises SEO ID NO: 9998.

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The invention also provides amino acid sequences having sequence identity to an amino acid sequence encoded by SEQ ID NO: 9968. The invention provides amino acids having sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO^S: 9969 to 10032. The degree of sequence identity is preferably greater than 50% (e.g., 60%, 70%, 80%, 85%, 88%, 90%, 92%, 95%, 99% or more).

A portion of SEQ ID NO: 9968 matches with approximately 98% identity to a previously published SARS polynucleotide sequence, commonly referred to as "BNI-1" (SEQ ID NO: 10033). BNI-1 was sequenced at Bernhard Nocht Institute for Tropical Medicine, National Reference Center for Tropical Infectious Diseases in Hamburg, Germany. The BNI-1 sequence was published on the WHO website on April 4, 2003 at http://www.who.int/csr/sars/primers/en and in Dorsten et al., "Identification of a Novel Coronavirus in Patients with Severe Acute Respiratory Syndrome", New England Journal of Medicine, published online at http://www.nejm.org on April 10, 2003. Both references are incorporated herein by reference in their entirety. The six reading frames of this 302mer sequence are shown in Figure 114 (see also Figure 129). The constituent amino acid sequences from Figure 114, having at least 4 amino acids, are listed as SEQ ID NOS: 10034 to 10065. An alignment of SEQ ID NO: 10034 with SEQ ID NO: 9997 is shown in Figure 130.

The invention provides for polynucleotide sequences comprising fragments of SEQ ID NO: 9968. In one embodiment, the fragment does not consist entirely of SEQ ID NO: 10033 or of a known coronavirus.

. The invention provides for amino acid sequences comprising fragments of an amino acid sequence encoded by SEQ ID NO: 9968. In one embodiment, the fragment does not consist entirely of an amino acid sequence encoded by SEQ ID NO: 10033 or a known coronavirus.

The invention provides for amino acids comprising fragments of an amino acid sequence selected from the group consisting of SEQ ID NO^S: 9969 to 10032. In one embodiment, the fragment does not consist entirely of an amino acid sequence encoded by the polynucleotide sequence of SEQ ID NO: 10033 or a known coronavirus.

Approximately 100 nucleotides at the 5' end of SEQ ID NO: 9968 do not match any portion of the BNI-1 polynucleotide sequence (SEQ ID NO: 10033). This unmatched portion is set forth as SEQ ID NO: 10066. The invention thus further provides a polynucleotide comprising the sequence comprising SEQ ID NO: 10066, polynucleotide sequences having sequence identity to SEQ ID NO: 10066, or polynucleotide sequences comprising fragments of SEQ ID NO: 10066.

The invention further comprises an amino acid sequence encoded by SEQ ID NO: 10066, an amino acid sequence having sequence identity to an amino acid sequence encoded by SEQ ID NO: 10066, or an amino acid sequence comprising fragments of an amino acid sequence

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encoded by SEQ ID NO: 10066. Preferably, the amino acid sequence comprises SEQ ID NO: 10067.

SEQ ID NO: 9997/9998 demonstrates homology with the a region of pollab of several coronaviruses. FIGURE 115 shows an alignment of SEQ ID NO^S: 9997/9998 to amino acid sequences for pollab of bovine coronavirus (SEQ ID NO: 10068), avian infectious bronchitis virus (SEQ ID NO: 10069) and murine hepatitis virus (SEQ ID NO: 10070). A consensus amino acid sequence of SEQ ID NO^S: 9997/9998, SEQ ID NO: 10068, SEQ ID NO: 10069, and SEQ ID NO: 10070 is shown in the bottom row of the alignment in Figure 115 (e.g. SEQ ID NO: 10071).

As shown in FIGURE 113, the polynucleotide sequence encoding SEQ ID NO: 9997 has a stop codon after codon 205, between SEQ ID NO^S: 9997 and 9998. Optionally, the stop codon can be removed and the amino acid sequence continued (SEQ ID NO: 10072). Accordingly, the invention provides for an amino acid sequence comprising SEQ ID NO: 9997 and/or SEQ ID NO: 9998, or SEQ ID NO: 10072, and further comprising an amino acid sequence encoding for the C-terminus of a coronavirus pollab gene or a fragment thereof.

As shown in FIGURE 115, SEQ ID NO^S: 10068, 10069, 10070 and 10071 contain amino acids prior to the N-terminus of SEQ ID NO: 9997. The invention also provides for an amino acid sequence comprising SEQ ID NO: 9997 and further comprising an amino acid sequence encoding for the N-terminus of a coronavirus pollab protein or a fragment thereof.

The pollab sequences on FIGURE 115 contain a coding region indicated on the schematic of FIGURE 117 by a "*". In FIGURE 115, the beginning of this genomic region is designated by the arrow crossing in front of amino acid 6080 of the consensus sequence SEQ ID NO: 10071. The end of this genomic region is designated by the arrow crossing in front of amino acid 6604 of the consensus sequence. The invention provides for an amino acid sequence comprising SEQ ID NO: 9997 and/or SEQ ID NO: 9998, or SEQ ID NO: 10072, and further comprising a first amino sequence prior to the N-terminus of said SEQ ID NO: 9997 and/or SEQ ID NO: 9998, or SEQ ID NO: 10072, wherein said first amino acid sequence has homology to an N-terminus sequence of a known coronavirus pollab "*" protein or a fragment thereof.

The invention further provides for an amino acid sequence comprising SEQ ID NO: 9997 and SEQ ID NO: 9998, wherein the stop codon after SEQ ID NO: 9971 is removed (i.e. SEQ ID NO: 10072), and further comprising a second amino acid sequence following the C terminus of SEQ ID NO: 9998, wherein said second amino acid sequence is homologous with a C terminus of a known coronavirus pollab "*" protein or a fragment thereof.

Examples of such proteins are shown aligned in FIGURE 118, and are SEQ ID NO⁵: 10073 to 10077. SEQ ID NO: 10073 comprises SEQ ID NO: 9997 and further comprises amino

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acids prior to the N-terminus and subsequent to the C-terminus from the pollab "*" protein of avian infectious bronchitis virus. SEQ ID NO: 10074 comprises SEQ ID NO: 9997 and further comprises amino acids prior to the N-terminus and subsequent to the C-terminus from the pollab "*" protein of bovine coronavirus. SEQ ID NO: 10075 comprises SEQ ID NO: 9997 and further comprises amino acids prior to the N-terminus and subsequent to the C-terminus from the pollab "*" protein of murine hepatitis virus. SEQ ID NO: 10076 comprises SEQ ID NO: 9997 and further comprises amino acids prior to the N-terminus and subsequent to the C-terminus from the consensus of the pollab "*" protein of avian infectious bronchitis virus, bovine coronavirus, and murine hepatitis virus (FIGURE 115). SEQ ID NO: 10077 comprises the consensus sequence of SEQ ID NOS: 10073 to 10076.

The invention comprises an amino acid sequence selected from the group consisting of SEQ ID NO^S: 10073, 10074, 10075, 10076 and 10077. The invention further includes an amino acid sequence comprising fragments of an amino acid sequence selected from the group consisting of SEQ ID NO^S: 10073, 10074, 10075, 10076 and 10077. The invention further comprises an amino acid sequence with sequence identity to a sequence selected from the group consisting of SEQ ID NO^S: 10073, 10074, 10075, 10076 and 10077.

The invention comprises polynucleotides encoding for the amino acid sequences selected from the group consisting of SEQ ID NO^S: 10073, 10074, 10075, 10076 and 10077. The invention comprises polynucleotides having sequence identity to polynucleotides encoding for the amino acid sequences selected from the group consisting of SEQ ID NO^S: 10073, 10074, 10075, 10076 and 10077. The invention comprises fragments of polynucleotides encoding SEQ ID NO^S: 10073, 10074, 10075, 10076 and 10077.

As shown in Figure 113, SEQ ID NO: 9968 includes a sequence that encodes SEQ ID NO: 10020 followed by a stop codon, giving a C-terminus threonine (Thr) residue. The corresponding sequence from an amino acid sequence encoded by BNI-1 is SEQ ID NO: 10078, which continues past the C-terminus of SEQ ID NO: 10020. Accordingly, the invention includes a protein comprising amino acid sequence SEQ ID NO: 10020 or an amino acid sequence having sequence identity to SEQ ID NO: 10020 or an amino acid sequence comprising a fragment of SEQ ID NO: 10020, wherein the C-terminus residue of said protein is a threonine. Preferably, the C-terminus of said protein is –ST. Still more preferably, the C-terminus of said protein is –ST. The invention also includes a protein comprising amino acid sequence SEQ ID NO: 10078 or an amino acid sequence having sequence identity to SEQ ID NO: 10078 or an amino acid sequence comprising a fragment of SEQ ID NO: 10078, wherein the C-terminus residue of said protein is Thr. Preferably, the C-terminus of said protein is –ST. Still more preferably, the C-terminus of said protein is –ST.

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SEQ ID NO: 9968 also encodes a 54mer amino acid sequence SEQ ID NO: 10015. The polynucleotide encoding SEQ ID NO: 10015 encodes two stop codons at its C-terminus (Figure 113). The corresponding region from the BNI-1 sequence does not contain this 54mer.

Accordingly, the invention includes a protein comprising amino acid sequence SEQ ID NO: 10015, or an amino acid sequence having sequence identity to SEQ ID NO: 10015 or an amino acid sequence comprising a fragment of SEQ ID NO: 10015. The invention further includes a polypeptide comprising SEQ ID NO: 10015 and further comprising a first amino acid sequence prior to the N-terminus of SEO ID NO: 10015.

SEQ ID NO: 9968 encodes the amino acid sequence SEQ ID NO: 9969. The polynucleotide sequence contains a stop codon at the C-terminus of SEQ ID NO: 9969. Accordingly, the invention includes a protein comprising amino acid sequence SEQ ID NO: 9969, or an amino acid sequence having sequence identity to SEQ ID NO: 9969. The invention further includes a polypeptide comprising SEQ ID NO: 9969 and further comprising a first amino acid sequence prior to the N-terminus of SEQ ID NO: 9969. The invention further includes a polypeptide comprising the sequence SEQ ID NO: 10079.

SEQ ID NO: 9968 encodes amino acid sequence QRT (Figure 113), followed by a stop codon. Accordingly, the invention includes a protein comprising amino acid sequence QRT. The invention further includes a polypeptide comprising amino acid sequence QRT and further comprising a first amino acid sequence prior to the N-terminus of the sequence QRT.

SEQ ID NO: 9968 encodes amino acid sequence SEQ ID NO: 10022, followed by a stop codon at its C-terminus. Accordingly, the invention includes a protein comprising amino acid sequence SEQ ID NO: 10022, or an amino acid sequence having sequence identity to SEQ ID NO: 10022. The invention further includes a polypeptide comprising SEQ ID NO: 10022 and further comprising a first amino acid sequence prior to the N-terminus of SEQ ID NO: 10022.

SEQ ID NO: 9968 encodes amino acid sequence SEQ ID NO: 10027. Within the SEQ ID NO: 10027 coding sequence there are at least three start codons, identified with underlining in Figure 119. The open reading frame indicated by the first start codon is SEQ ID NO: 10081. The open reading frame indicated by the second start codon is SEQ ID NO: 10082. The open reading frame indicated by the third start codon is SEQ ID NO: 10083.

The invention provides a novel SARS polynucleotide sequence SEQ ID NO: 10084. All six reading frames of this 1463mer sequence are shown in Figure 120 (see also Figure 122). The constituent amino acid sequences from Figure 120, having at least 4 amino acids, are listed as SEQ ID NOS: 10085 to 10209 (see Figures 120A to 120F).

The invention includes a polynucleotide sequence comprising SEQ ID NO: 10084. The invention also provides polynucleotide sequences having sequence identity to SEQ ID NO: 10084. The invention also provides for polynucleotide sequences comprising fragments of SEQ

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ID NO: 10084. In one embodiment, the polynucleotide fragment does not consist entirely of SEQ ID NO: 10033 or a known coronavirus polynucleotide sequence or a known SARS polynucleotide sequence.

The invention includes an amino acid sequence encoded by the polynucleotide sequence of SEQ ID NO: 10084, including the amino acid sequences of Figures 120A to 120F e.g. selected from the group consisting of SEQ ID NO^S: 10085 to 10209. Preferably, the amino acid sequence comprises SEQ ID NO: 10149.

The invention also provides amino acid sequences having sequence identity to an amino acid sequence encoded by SEQ ID NO: 10084. The invention provides amino acids having sequence identity to an amino acid sequence from Figures 120A to 120F e.g. selected from the group consisting of SEQ ID NO^S: 10085 to 10209.

The invention also provides fragments of amino acid sequences encoded by SEQ ID NO: 10084. The invention also provides fragments of amino acid sequences selected from the group consisting of SEQ ID NO^S: 10085 to 10209. In one embodiment, the fragment does not consist entirely of an amino acid sequence encoded by SEQ ID NO: 10033 or an amino acid sequence of a known coronavirus or an amino acid sequence of a known SARS virus. An alignment of the matching portion of SEQ ID NO: 10033 and SEQ ID NO: 10084 is included in FIGURE 121.

In one embodiment, the invention comprises an amino acid sequence comprising SEQ ID NO: 10149. An alignment of the polynucleotide sequence SEQ ID NO: 10084 to the encoded SEQ ID NO: 10149 is shown in FIGURE 122 (5'3' Frame 3). Analysis of the 5'3' Frame 3 translation by a computer program to predict start codon methionines (NetStart 1.0) (FIGURE 123) reveals SEQ ID NO^S: 10210 to 10215.

The invention includes a protein comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10210, SEQ ID NO: 10211, SEQ ID NO: 10212, SEQ ID NO: 10213, SEQ ID NO: 10214 and SEQ ID NO: 10215. The invention includes a protein having sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO: 10210, SEQ ID NO: 10211, SEQ ID NO: 10212, SEQ ID NO: 10213, SEQ ID NO: 10214 and SEQ ID NO: 10215. In one embodiment, the protein does not consist entirely of an amino acid sequence of a known SARS virus or of a known coronavirus.

The invention includes a fragment of a protein comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10210, SEQ ID NO: 10211, SEQ ID NO: 10212, SEQ ID NO: 10213, SEQ ID NO: 10214 and SEQ ID NO: 10215. In one embodiment, the fragment does not consist entirely of an amino acid sequence of a known SARS virus or of a known coronavirus.

In one embodiment, the invention includes a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 10210, SEQ ID NO: 10211 and

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SEQ ID NO: 10212. Partial results of a BLAST of SEQ ID NO: 10210 against GenBank is included in FIGURE 124. These results indicate that SEQ ID NOS: 10210, 10211 and 10212 have functional similarities to a Coronavirus RNA polymerase, particularly the RNA polymerase of murine hepatitis virus, bovine coronavirus, and avian infectious bronchitis.

In one embodiment, the invention is directed to a polypeptide comprising a first amino acid sequence selected from the group consisting of SEQ ID NO: 10210, SEQ ID NO: 10211 and SEQ ID NO: 10212 and a second amino acid sequence from the C-terminus of a coronavirus ORF1ab sequence. Preferably, the second amino acid sequence is from a bovine coronavirus. One example of this embodiment is shown below as SEQ ID NO: 10216. Amino acids 1-481 of SEQ ID NO: 10216 are the first amino acid sequence of SEQ ID NO: 10210, and amino acids 482-1152 are the second amino acid sequence of the C-terminus of a bovine coronavirus or flab polyprotein (Gi 26008080) (NP_150073.2) (SEQ ID NO: 10217).

Accordingly, the invention includes a polypeptide comprising SEQ ID NO: 10216. The invention further includes a polypeptide comprising a first amino acid sequence of SEQ ID NO: 10210 and a second amino acid sequence of SEQ ID NO: 10217. The invention further includes a polypeptide comprising a first amino acid sequence having greater than x% identity to SEQ ID NO: 10210 and a second amino acid sequence having greater than y% identity to SEQ ID NO: 10217, wherein x is greater than or equal to 85% (e.g., 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more) and wherein y is greater than or equal to 60% (e.g., 65%, 70%, 75%, 80%, 85%, 90%, 95%, 98%, 99% or more).

The invention also includes a polypeptide comprising a fragment of SEQ ID NO: 10210, wherein said fragment includes an epitope. Computer-predicted epitopes of SEQ ID NO: 10210, using a 17mer window, are included in FIGURE 125A (Hopp & Woods) and FIGURE 125B (Kyte & Doolittle).

The amino acid sequence of SEQ ID NO: 10210 also contains two predicted glycosylation sites at amino acids 81–84 (NNTE; SEQ ID NO: 10218) and at 180–183 (NHSV; SEQ ID NO: 10219). Accordingly, the invention includes a polypeptide comprising a fragment of SEQ ID NO: 10210, wherein said fragment includes a glycosylation site. The invention further includes a polypeptide comprising a fragment of SEQ ID NO: 10210, wherein said fragment includes the Asn at position 81. Preferably, said Asn is glycosylated. The invention further includes a polypeptide comprising a fragment of SEQ ID NO: 10210, wherein said fragment includes the Asn at position 180. Preferably, said Asn is glycosylated.

In one embodiment, the invention includes a polypeptide comprising an amino acid sequence from within Figure 120D and/or SEQ ID NO^S: 10150 to 10160 e.g. from SEQ ID NO^S: 10154, 10155, 10158 and 10160. Within SEQ ID NO: 10154 the following amino acid sequences starting with a Met and ending at a stop codon can be identified: SEQ ID NO^S: 10220 to 10227.

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Accordingly, the invention includes a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10220, SEQ ID NO: 10221, SEQ ID NO: 10222, SEQ ID NO: 10223, SEQ ID NO: 10224, SEQ ID NO: 10225, SEQ ID NO: 10226 and SEQ ID NO: 10227, or a fragment thereof or an amino acid sequence having sequence identity thereto.

In one embodiment, the invention includes a polypeptide comprising the amino acid sequence within Figure 120E e.g. from SEQ ID NOS: 10161 to 10182, and in particular SEQ ID NOS: 10171 and 10176. Within SEQ ID NOS: 10171 and 10176 the following amino acid sequences starting with a Met and ending at a stop codon can be identified: SEQ ID NO: 10228 and SEQ ID NO: 10229.

Accordingly, the invention includes a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10228 and SEQ ID NO: 10229, or a fragment thereof or an amino acid sequence having sequence identity thereto.

In one embodiment, the invention includes a polypeptide comprising an amino acid sequence from Figure 120F e.g. SEQ ID NO^S: 10183 to 10209. Within Figure 120F the following amino acid sequence starting with a Met and ending at a stop codon can be identified: SEQ ID NO: 10187. Accordingly, the invention includes a polypeptide comprising an amino acid sequence of SEQ ID NO: 10187, or a fragment thereof or an amino acid sequence having sequence identity thereto.

In one embodiment, the polynucleotides of the invention do not include one of the following primers, disclosed at http://content.nejm.org/cgi/reprint/NEJMoa030781v2.pdf;

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5'GGGTTGGGACTATCCTAAGTGTGA3' (SEQ ID NO: 10230)
5'TAACACACACTCCATCCATCA3' (SEQ ID NO: 10231)
5'CTAACATGCTTAGGATAATGG3' (SEQ ID NO: 10232)
5'GCCTCTCTTGTTCTTGCTCGC3' (SEQ ID NO: 10234)
5'CAGGTAAGCGTAAAACTCATC3' (SEQ ID NO: 10234)
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The invention also includes polynucleotide sequences which can be used as probes for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes the polynucleotide primers identified in Table 31 (SEQ ID NO⁸: 10235 to 10258), the forward primers SEQ ID NO⁸: 10259 to 10281 and the reverse primers SEQ ID NO⁸: 10282 to 10298. The invention further includes polynucleotide sequences which are complementary to any one of these primer sequences disclosed herein.

The invention provides a SARS polynucleotide sequence SEQ ID NO: 10299. All six reading frames of this sequence are included in FIGURE 126 (See also Figure 131). The constituent amino acid sequences from Figure 126, having at least 4 amino acids, are listed as SEQ ID NOS: 10300 to 10337.

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Accordingly, the invention includes a polynucleotide sequence comprising SEQ ID NO: 10299. It also provides polynucleotide sequences having sequence identity to SEQ ID NO: 10299. The invention also provides for polynucleotide sequences comprising fragments of SEQ ID NO: 10299. In one embodiment, the polynucleotide fragment does not consist entirely of a known polynucleotide sequence of a SARS virus or a known polynucleotide sequence of a coronavirus.

The invention includes an amino acid sequence encoded by the polynucleotide sequence of SEQ ID NO: 10299, including the amino acid sequences shown in Figure 126, and the amino acid sequences selected from the group consisting of SEQ ID NO^S: 10300 to 10337. Preferably, the amino acid sequence comprises SEO ID NO: 10316.

The invention also provides amino acid sequences having sequence identity to an amino acid sequence encoded by SEQ ID NO: 10299. The invention provides amino acid sequences having identity to an amino acid sequence selected from the group consisting of SEQ ID NO^S: 10300 to 10337.

The invention also provides fragments of amino acid sequences encoded by SEQ ID NO: 10299. The invention also provides fragments of amino acid sequences selected from the group consisting of SEQ ID NO^S: 10300 to 10337. In one embodiment, the fragment does not consist entirely of a known amino acid sequence of a SARS virus or a known amino acid sequence of a coronavirus.

In one embodiment, the invention comprises an amino acid sequence comprising SEQ ID NO: 10316. Encoded open reading frames within SEQ ID NO: 10316 include SEQ ID NO: 10338 and SEO ID NO: 10339.

In one embodiment, the invention comprises an amino acid sequence comprising a sequence from within the 5'3' Frame 1 translation of SEQ ID NO: 10299. The following encoded open reading frame is found within this translation: SEQ ID NO: 10340.

In one embodiment, the invention comprises an amino acid sequence comprising a sequence from within the 3'5' Frame 1 translation of SEQ ID NO: 10299. An encoded open reading frame within this translation is SEQ ID NO: 10341.

In one embodiment, the invention comprises an amino acid sequence comprising a sequence from within the 3'5' Frame 2 translation of SEQ ID NO: 10299. An encoded open reading frame within this translation is SEQ ID NO: 10342.

The invention includes a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10338, SEQ ID NO: 10339, SEQ ID NO: 10340, SEQ ID NO: 10341 and SEQ ID NO: 10342. The invention includes a polypeptide having sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO: 10338, SEQ ID NO: 10340, SEQ ID NO: 10341 and SEQ ID NO: 10342. The invention

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includes a fragment of a polypeptide comprising an amino acid sequence elected from the group consisting of SEQ ID NO: 10338, SEQ ID NO: 10339, SEQ ID NO: 10340, SEQ ID NO: 10341 and SEQ ID NO: 10342. In one embodiment, the fragment does not consist entirely of a known SARS virus amino acid sequence or of a known coronavirus amino acid sequence.

In one embodiment, SEQ ID NOS: 10338-10342 are used in fusion proteins. Accordingly, the start codon methionines may be removed. The invention comprises a amino acid sequence selected from the group consisting of SEQ ID NO: 10343, SEQ ID NO: 10344, SEQ ID NO: 10345, SEQ ID NO: 10346 and SEQ ID NO: 10347.

In one embodiment, the invention comprises an amino acid sequence selected from the group consisting of SEQ ID NO: 10338 and SEQ ID NO: 10339. Partial BLAST results of SEQ ID NO: 10338 against GenBank are given below:

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>gi|133593|sp|P18457|RRPB_CVPFS RNA-DIRECTED RNA POLYMERASE (ORF1B)
gi|93934|pir||A43489 RNA-directed RNA polymerase (EC 2.7.7.48) - porcine
transmissible gastroenteritis virus (fragment)
gi|833161|emb|CA37284.1| polymerase [Transmissible gastroenteritis:
virus]
Length = 533
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Score = 131 bits (329), Expect = 3e-30

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BNSDOCID: <WO ____ 2004092360A2 | >

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Identities = 55/89 (61%), Positives = 69/89 (77%), Gaps = 1/89 (1%)
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Query: 1 MLWCKDGHVETFYPKLQASQAWQPGVAMPNLYKMQRMLLEKCDLQNYGENAVIPKGIMMN 60 MLWC++ H++TFYP+LQ+++ W PG +MP LYX+QRM LE+C+L NYG +P GI N Sbjct: 217 MLWCENSHIKTFYPQLQSAE-WNPGYSMPTLYKIQRMCLERCNLYNYGAQVKLPDGITTM 275

Query: 61 VAKYTOLCQYLNTLTLAVPSNMRVIHFGA 89
V KYTQLCQYLNT TL VP MRV+H GA
Sbjet: 276 VVKYTQLCQYLNTTTLCVPHKMRVLHLGA 304

These results indicate that SEQ ID NO: 10338 has functional similarities to an RNA-directed RNA polymerase of porcine transmissible gastroenteritis virus.

Partial BLAST results of SEQ ID NO: 10339 against GenBank are given below: >gb|AAL57305.1| replicase [bovine coronavirus]

Length = 7094

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Score = 139 bits (351), Expect = 7e-33
Identities = 64/108 (59%); Positives = 78/108 (72%)
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Query: 1 MSVISKVVKVTIDYAEISFMLWCKDGHVETFYPKLQASQAWQPGVAMPNLYKMQRMLLEK 60
M+ +SKVV V +D+ + FMLWC D V TFYP+LQA+ W+PG +MP LYK +E+
55jct: 6760 LNCVSKVVNVNVDFKDFQFMLWCNDEKVMTFYPRLQAASDWKPGYSMPVLYKYLNSPMER

Query: 61 CDLQNYGENAVIPKGIMMNVAKYTQLCQYLNTLTLAVPSNMRVIHFGA 108 L NYG+ +P G MMNVAKYTQLCQYLNT TLAVP NMRV+H GA Sbjct: 6820 VSLWNYGKPVTLPTGCMMNVAKYTQLCQYLNTTTLAVPVNMRVLHLGA 6867

These results indicate that SEQ ID NO: 10339 has functional similarities to a replicase of bovine coronavirus.

The SARS virus may contain polymorphism at the Glu-20 residue of SEQ ID NO: 10338. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 10338, wherein said polypeptide includes an amino acid sequence selected from the group consisting of ASQAW (SEQ ID NO: 10348) and ASRAW (SEQ ID NO: 10349). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 10338, wherein said fragment includes an amino acid sequence selected from the group consisting of SEO ID NO: 10348 and SEO ID NO: 10349.

The SARS virus may contain polymorphism at the Ser-80 residue of SEQ ID NO: 10338. below. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 10338, wherein said polypeptide includes an amino acid sequence selected from the group consisting of VPSNM (SEQ ID NO: 10350) and VPTNM (SEQ ID NO: 10351). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 10338, wherein said fragment includes an amino acid sequence selected from the group consisting of SEQ ID NO: 10350 and SEQ ID NO: 10351.

The invention also includes polynucleotide sequences which can be used as probes for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in Table 32. The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in Table 32.

The invention provides a SARS polynucleotide sequence SEQ ID NO: 10505. All six reading frames of this sequence are shown in Figure 127 (see also Figure 132). The constituent amino acid sequences from Figure 127, having at least 4 amino acids, are listed as SEQ ID NOS: 10506 to 10570.

The invention includes a polynucleotide sequence comprising SEQ ID NO: 10505. The invention also provides polynucleotide sequences having sequence identity to SEQ ID NO: 10505. The invention also provides for polynucleotide sequences comprising fragments of SEQ ID NO: 10505. In one embodiment, the polynucleotide fragment does not consist entirely of a known SARS virus polynucleotide sequence or of a known coronavirus polynucleotide sequence.

The invention includes an amino acid sequence encoded by the polynucleotide sequence of SEQ ID NO: 10505, including the amino acid sequences shown in Figure 127, and particularly those selected from the group consisting of SEQ ID NO^S: 10506 to 10570. Preferably, the amino acid sequence comprises SEQ ID NO: 10532 and/or SEQ ID NO: 10533.

The invention also provides amino acid sequences having sequence identity to an amino acid sequence encoded by SEQ ID NO: 10505. The invention provides amino acid sequences

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As used herein, the term "target nucleic acid region" or "target nucleic acid" denotes a nucleic acid molecule with a "target sequence" to be amplified. The target nucleic acid may be either single-stranded or double-stranded and may include other sequences besides the target sequence, which may not be amplified. The term "target sequence" refers to the particular nucleotide sequence of the target nucleic acid which is to be amplified. The target sequence may include a probe-hybridizing region contained within the target molecule with which a probe will form a stable hybrid under desired conditions. The "target sequence" may also include the complexing sequences to which the oligonucleotide primers complex and be extended using the target sequence as a template. Where the target nucleic acid is originally single-stranded, the term "target sequence" also refers to the sequence complementary to the "target sequence" as present in the target nucleic acid. If the "target nucleic acid" is originally double-stranded, the term "target sequence" refers to both the plus (+) and minus (-) strands.

The term "primer" or "oligonucleotide primer" as used herein, refers to an oligonucleotide which acts to initiate synthesis of a complementary DNA strand when placed under conditions in which synthesis of a primer extension product is induced *i.e.* in the presence of nucleotides and a polymerization-inducing agent such as a DNA or RNA polymerase and at suitable temperature, pH, metal concentration, and salt concentration. The primer is preferably single-stranded for maximum efficiency in amplification, but may alternatively be double-stranded. If double-stranded, the primer is first treated to separate its strands before being used to prepare extension products. This denaturation step is typically effected by heat, but may alternatively be carried out using alkali, followed by neutralization. Thus, a "primer" is complementary to a template, and complexes by hydrogen bonding or hybridization with the template to give a primer/template complex for initiation of synthesis by a polymerase, which is extended by the addition of covalently bonded bases linked at its 3' end complementary to the template in the process of DNA synthesis.

As used herein, the term "probe" or "oligonucleotide probe" refers to a structure comprised of a polynucleotide, as defined above, that contains a nucleic acid sequence complementary to a nucleic acid sequence present in the target nucleic acid analyte. The polynucleotide regions of probes may be composed of DNA, and/or RNA, and/or synthetic nucleotide analogs. When an "oligonucleotide probe" is to be used in a 5' nuclease assay, such as the TaqManTM technique, the probe will contain at least one fluorescer and at least one quencher which is digested by the 5' endonuclease activity of a polymerase used in the reaction in order to detect any amplified target oligonucleotide sequences. In this context, the oligonucleotide probe will have a sufficient number of phosphodiester linkages adjacent to its 5' end so that the 5' to 3' nuclease activity employed can efficiently degrade the bound probe to separate the fluorescers and quenchers.

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When an oligonucleotide probe is used in the TMA technique, it will be suitably labeled, as described below.

It will be appreciated that the hybridizing sequences need not have perfect complementarity to provide stable hybrids. In many situations, stable hybrids will form where fewer than about 10% of the bases are mismatches, ignoring loops of four or more nucleotides. Accordingly, as used herein the term "complementary" refers to an oligonucleotide that forms a stable duplex with its "complement" under assay conditions, generally where there is about 90% or greater homology.

The terms "hybridize" and "hybridization" refer to the formation of complexes between nucleotide sequences which are sufficiently complementary to form complexes via Watson-Crick base pairing. Where a primer "hybridizes" with target (template), such complexes (or hybrids) are sufficiently stable to serve the priming function required by e.g. the DNA polymerase to initiate DNA synthesis.

Stringent hybridization conditions will typically include salt concentrations of less than about 1 M, more usually less than about 500 mM and preferably less than about 200 mM. Hybridization temperatures can be as low as 5°C, but are typically greater than 22°C, more typically greater than about 30°C, and preferably in excess of about 37°C. Longer fragments may require higher hybridization temperatures for specific hybridization. Other factors may affect the stringency of hybridization, including base composition and length of the complementary strands, presence of organic solvents and extent of base mismatching, and the combination of parameters used is more important than the absolute measure of any one alone. Other hybridization conditions which may be controlled include buffer type and concentration. solution pH, presence and concentration of blocking reagents to decrease background binding such as repeat sequences or blocking protein solutions, detergent type(s) and concentrations. molecules such as polymers which increase the relative concentration of the polynucleotides, metal ion(s) and their concentration(s), chelator(s) and their concentrations, and other conditions known in the art. Less stringent, and/or more physiological, hybridization conditions are used where a labeled polynucleotide amplification product cycles on and off a substrate linked to a complementary probe polynucleotide during a real-time assay which is monitored during PCR amplification such as a molecular beacon assay. Such less stringent hybridization conditions can also comprise solution conditions effective for other aspects of the method, for example reverse transcription or PCR.

As used herein, a "biological sample" refers to a sample of tissue, cells or fluid isolated from a subject, that commonly includes antibodies produced by the subject. Typical samples include but are not limited to, blood, plasma, serum, fecal matter, urine, bone marrow, bile, spinal fluid, lymph fluid, samples of the skin, secretions of the skin, respiratory, intestinal, and

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genitourinary tracts, tears, saliva, sputum, mucous, milk, blood cells, organs, tissues, biopsies (e.g. lung, liver, kidney) and also samples of in vitro cell culture constituents including but not limited to conditioned media resulting from the growth of cells and tissues in culture medium e.g. recombinant cells, and cell components. Other samples that may be used for diagnosis include stool samples and nasopharyngeal aspirates.

The term "antibody" encompasses polyclonal and monoclonal antibody preparations, as well as preparations including hybrid antibodies, altered antibodies, chimeric antibodies and, humanized antibodies, as well as: hybrid (chimeric) antibody molecules (see, for example, Winter et al. (1991) Nature 349:293-299; and US Patent 4,816,567); F(ab')₂ and F(ab) fragments; Fv molecules (noncovalent heterodimers, see, for example, Inbar et al. (1972) Proc Natl Acad Sci USA 69:2659-2662; and Ehrlich et al. (1980) Biochem 19:4091-4096); single-chain Fv molecules (sFv) (see, e.g., Huston et al. (1988) Proc Natl Acad Sci USA 85:5879-5883); oligobodies; dimeric and trimeric antibody fragment constructs; minibodies (see, e.g., Pack et al. (1992) Biochem 31:1579-1584; Cumber et al. (1992) J Immunology 149B:120-126); humanized antibody molecules (see, e.g., Riechmann et al. (1988) Nature 332:323-327; Verhoeyan et al. (1988) Science 239:1534-1536; and UK Patent Publication No. GB 2,276,169, published 21 September 1994); and, any functional fragments obtained from such molecules, wherein such fragments retain specific-binding properties of the parent antibody

As used herein, the term "monoclonal antibody" refers to an antibody composition having a homogeneous antibody population. The term is not limited regarding the species or source of the antibody, nor is it intended to be limited by the manner in which it is made. The term encompasses whole immunoglobulins.

Methods of making polyclonal and monoclonal antibodies are known in the art. Polyclonal antibodies are generated by immunizing a suitable animal, such as a mouse, rat, rabbit, sheep or goat, with an antigen of interest. In order to enhance immunogenicity, the antigen can be linked to a carrier prior to immunization. Suitable carriers are typically large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, lipid aggregates (such as oil droplets or liposomes), and inactive virus particles. Such carriers are well known to those of ordinary skill in the art. Furthermore, the antigen may be conjugated to a bacterial toxoid, such as toxoid from diphtheria, tetanus, cholera, etc., in order to enhance the immunogenicity thereof.

Rabbits, sheep and goats are preferred for the preparation of polyclonal sera when large volumes of sera are desired. These animals are good design choices also because of the availability of labeled anti-rabbit, anti-sheep and anti-goat antibodies. Immunization is generally performed by mixing or emulsifying the antigen in saline, preferably in an adjuvant such as

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Freund's complete adjuvant ("FCA"), and injecting the mixture or emulsion parenterally (generally subcutaneously or intramuscularly). The animal is generally boosted 2-6 weeks later with one or more injections of the antigen in saline, preferably using Freund's incomplete adjuvant ("FIA"). Antibodies may also be generated by in vitro immunization, using methods known in the art. Polyclonal antisera is then obtained from the immunized animal.

Monoclonal antibodies are generally prepared using the method of Kohler & Milstein (1975) Nature 256:495-497, or a modification thereof, as described above.

Nucleic acid detection methods

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There are many well known methods of amplifying targeted sequences, such as the polymerase chain reaction (PCR), reverse transcription PCR (RT-PCR), the ligase chain reaction (LCR), the strand displacement amplification (SDA), and the nucleic acid sequence-based amplification (NASBA), transcription-mediated amplification (TMA) to name a few. These methods are described generally in the following references: (PCR) US Patents 4,683,195, 4,683,202, and 4,800,159; (RT-PCR) US patent 5,310,652, 5,322,770; (LCR) EP Application No., 320,308 published Jun. 14, 1989; (SDA) US Pat. Nos. 5,270,184, and 5,455,166 and "Empirical Aspects of Strand Displacement Amplification" by G. T. Walker in PCR Methods and Applications, 3(1):1-6 (1993), Cold Spring Harbor Laboratory Press; (TMA) US Patent No. 5,399,491, and (NASBA) "Nucleic Acid Sequence-Based Amplification (NASBATM)" by L. Malek et al., Ch. 36 in Methods in Molecular Biology, Vol. 28: Protocols for Nucleic Acid

Analysis by Nonradioactive Probes, 1994 Ed. P. G. Isaac, Humana Press, Inc., Totowa, N.J. PCR methods may include variations that permit quantitation of the target sequence, for example, by real time PCR analysis (e.g., as described in US patents 5,210,015, 5,487,972, 5,994,056, 6,171,785 inter alia). (Each of the above references are hereby incorporated by reference).

One embodiment of the method of the invention for detecting the presence of SARS virus in a sample comprises providing a sample suspected of containing a SARS virus nucleic acid target, amplifying a template sequence contained within said SARS virus nucleic acid target by any known technique of nucleic acid amplification, including any of those mentioned herein, using the oligonucleotide primers described herein, particularly those primers comprising the kits described herein, and detecting the amplified template sequence, wherein the presence of the amplified template sequence indicates the presence of SARS virus in said sample.

Amplification techniques generally involve the use of two primers. Where a target sequence is single-stranded, the techniques generally involve a preliminary step in which a complementary strand is made in order to give a double-stranded target. The two primers hybridize to different strands of the double-stranded target and are then extended. The extended products can serve as targets for further rounds of hybridization/extension. The net effect is to amplify a template sequence within the target, the 5' and 3' termini of the template being defined

by the locations of the two primers in the target. As an alternative, if one or both of the primers contains a promoter sequence then the target can be amplified (by transcription) using a RNA polymerase (as in TMA).

The present invention provides methods and kits for amplifying and/or detecting a template or target sequence in the SARSV viral nucleic acid. The invention provides a kit comprising primers for amplifying a template sequence contained within a SARSV nucleic acid target, the kit comprising a first primer and a second primer, wherein the first primer comprises a sequence substantially complementary to a portion of said template sequence and the second primer comprises a sequence substantially complementary to a portion of the complement of said template sequence, wherein the sequences within said primers which have substantial complementarity define the termini of the template sequence to be amplified.

Kits of the invention may further comprise a probe which is substantially complementary to the template sequence and/or to its complement and which can hybridize thereto. This probe can be used in a hybridization technique to detect amplified template, or to isolate (i.e. "capture) the amplified template or the original target nucleic acid.

Kits of the invention may further comprise primers and/or probes for generating and detecting an internal standard, in order to aid quantitative measurements (e.g Fille et al. 1997 Biotechniques 23:34-36).

Kits of the invention may further comprise a DNA polymerase, which will generally be a thermostable DNA polymerase where a non-isothermal amplification process is to be used. The kits may also comprise supplies of dNTPs, a magnesium salt (e.g. MgCl₂), buffer solutions, etc.

Kits of the invention may comprise more than one pair of primers (e.g. for nested amplification), and one primer may be common to more than one primer pair. The kit may also comprise more than one probe.

Oligomer Probes and Primers

In connection with the nucleic acid detection methods of the present invention described above, oligomers having sequence similarity, or complementarity, to the SARSV genome are useful. The SARSV genome sequences mentioned herein may be used to produce probes and primers which can be used in assays for the detection of nucleic acids in test samples. The probes may be designed from conserved nucleotide regions of the polynucleotides of interest or from non-conserved nucleotide regions of the polynucleotide of interest. The design of such probes for optimization in assays is within the skill of those of ordinary skill in the art. Generally, nucleic acid probes are developed from non-conserved or unique regions when maximum specificity is desired, and nucleic acid probes are developed from conserved regions when assaying for nucleotide regions that are closely related to, for example, different members of a multi-gene family or in related species like mouse and man.

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Using as a basis the SARSV genome which can be found as described herein, and/or preferably conserved regions of the SARSV genome, and/or the particularly described primer and probe sequences as disclosed herein, oligomers of approximately 8 nucleotides or more can be prepared which hybridize with the positive strand(s) of SARSV RNA or its complement, as well as to SARSV cDNAs. These oligomers can serve as probes for the detection (including isolation and/or labeling) of polynucleotides which contain SARSV nucleotide sequences, and/or as primers for the transcription and/or replication of targeted SARSV sequences. The oligomers contain a targeting polynucleotide sequence, which is comprised of nucleotides which are complementary to a target SARSV nucleotide sequence; the sequence is of sufficient length and complementarity with the SARSV sequence to form a duplex which has sufficient stability for the purpose intended. For example, if the purpose is the isolation, via immobilization, of an analyte containing a target SARSV sequence, the oligomers would contain a polynucleotide region which is of sufficient length and complementarity to the targeted SARSV sequence to afford sufficient duplex stability to immobilize the analyte on a solid surface, via its binding to the oligomers, under the isolation conditions. For example, also, if the oligomers are to serve as primers for the transcription and/or replication of target SARSV sequences in an analyte polynucleotide, the oligomers would contain a polynucleotide region of sufficient length and complementarity to the targeted SARSV sequence to allow the polymerizing agent to continue replication from the primers which are in stable duplex form with the target sequence, under the polymerizing conditions. For example, also, if the oligomers are to be used as label probes, or are to bind to multimers, the targeting polynucleotide region would be of sufficient length and complementarity to form stable hybrid duplex structures with the label probes and/or multimers to allow detection of the duplex. The oligomers may contain a minimum of about 4 contiguous nucleotides which are complementary to targeted SARSV sequence; usually the oligomers will contain a minimum of about 8 contiguous nucleotides which are complementary to the targeted SARSV sequence, and preferably will contain a minimum of about 14, 15, 16, 17, 18, 19, 20,21, 22, 23, 24, 25, 26, 27, 28, 29, or 30 contiguous nucleotides and up to about 50, 75, 100, 200 contiguous nucleotides or more, which are complementary to the targeted SARSV sequence.

Typically, for use in the amplification based methods (for example, PCR, RT-PCR, TMA) oligomers will be used as primer sets such that one member of the primer set has sequence similarity or complementarity to a more conserved (among coronaviruses) portion of the SARSV genome and the other member of the primer set has sequence similarity or complementarity to a less conserved portion. The primer sets can be used to amplify the target region in ways that are well known in the art. Typically, the 5' untranslated region (5'UTR) and the 3' untranslated region (3'UTR) are among the most conserved regions. Figure 8 shows an alignment of the 5'UTR of several

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coronaviruses. Figures 9 and 11 show the sequences of preferred primers for amplification of the 5'UTR and 3'UTR, respectively. Other primers and probes can readily be designed based on the sequence alignments provided herein.

The oligomer, however, need not consist only of the sequence which is complementary to the targeted SARSV sequence. It may contain in addition, nucleotide sequences (e.g. promoters) or other moieties which are suitable for the purposes for which the oligomers are used. For example, if the oligomers are used as primers for the amplification of SARSV sequences via, for example, PCR, they may contain sequences which, when in duplex, form restriction enzyme sites which facilitate the cloning of the amplified sequences. For example, also, if the oligomers are to be used as "capture probes" in hybridization assays, they would contain in addition a binding partner which is coupled to the oligomer containing the nucleotide sequence which is complementary to the targeted SARSV sequence. Other types of moietites or sequences which are useful of which the oligomers may be comprised or coupled to, are those which are known in the art to be suitable for a variety of purposes, including the labeling of nucleotide probes.

Table 4 (SEQ ID NOS: 1021-6020) shows forward and reverse primers that are useful for nucleic acid amplification of SARSV for diagnostic and screening methods.

Preferred primers and probes for SARS nucleic acid detection for diagnostic and screening are SEQ ID NOS: 7332-7336 (forward primers), SEQ ID NOS: 7337-7341 (reverse primers) and SEQ ID NOS: 7342-7352 (probes). These primers and probes are useful for detection of sequences in the 3' UTR.

Any of the above forward primers may be used in combination with any of the above reverse primers for amplification of SARSV nucleic acid. The amplified product may be detected (or captured) with any of the above probes. Particularly preferred combinations of forward and reverse primers and the probes for detecting the amplified product include: Forward SEQ ID NO: 7332 with reverse SEQ ID NO: 7337, 7338, 7339 or 7341 and probe SEQ ID NO: 7342; forward SEQ ID NO: 7333 or 7334 with reverse SEQ ID NO: 7340 and any of probes SEQ ID NO: 7343-7351; Forward SEQ ID NO: 7335 and reverse SEQ ID NO: 7340 or 7341 and any of probes SEQ ID NO: 7342-7352. Other combinations of forward and reverse primers and appropriate probes can readily be determined by those skilled in the art from the above information.

Additional preferred primers and probes for SARS nucleic acid detection for diagnostic and screening are SEQ ID NOS: 7353-7362 (forward primers), SEQ ID NOS: 7363-7373 (reverse primers) and SEQ ID NOS: 7374-7385 (probes). The primers and probes are useful for detection of sequences in the 5' UTR.

The above primers may be used in combination for amplification of SARSV nucleic acid as follows: any of forward primers SEQ ID NO: 7353-7356 with any of reverse primers SEQ ID

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NO: 7363-7366, 7368 and the amplified product detected (or captured) with probes SEQ ID NO: 7374; any of forward primers SEQ ID NO: 7357-7362 with any of reverse primers SEQ ID NO: 7367, 7369-7373 and the amplified products detected (or captured) with any of probes SEO ID NO: 7375-7385. Particularly preferred combinations of forward and reverse primers and probes are: Forward primers SEO ID NO: 7353-7356 with any of reverse primers SEO ID NO: 7363-5 7366 and probes SEQ ID NO: 7374; forward primers SEO ID NO: 7357-7358 with reverse primers SEO ID NO: 7367, 7369 and probes SEO ID NO: 7375 or 7376; Forward primers SEO ID NO: 7357-7359 with reverse primers SEO ID NO: 7367, 7369 or 7370 and probe SEQ ID NO: 7375 or 7376. More preferred are combinations of SEQ ID NO: 7353 or 7354 with SEQ ID NO: 7363 or 7364 and probe SEO ID NO: 7374. Other combinations of forward and reverse primers and appropriate probes can readily be determined by those skilled in the art from the above information. A particularly conserved octanucleotide sequence (SEQ ID NO: 7386) occurs in the 3'UTR of SARS (approximately 70-80 bases from the 3' end) and of several other Coronaviruses that may be particularly useful in identifying SARSV. Primers including in this region are preferably combined with reverse primers from regions of sequence that are more specific for SARS.

In addition to the above, the intergenic sequence (IS) that is characteristic of Coronavirus has been identified in SARSV (see above). The IS minimally comprises the sequence ACGAAC (SEQ ID NO: 7293) which occurs upstream of each open reading frame (ORF) in the viral genome. The 5'UTR which includes the IS is spliced onto the 5' end of each viral mRNA at or adjacent to the site of the IS. Thus, primers comprising the IS or its complement are useful for amplifying viral nucleic acids, including cDNA made from the viral RNAs. The invention thus comprises a set of primers in which one primer comprises ACGAAC (SEQ ID NO: 7293) or its complement (SEQ ID NO: 7387) and one primer comprises any appropriate sequence from the SARS genome, or a complementary sequence. Useful probes for detecting and/or capturing the viral RNAs or cDNA made from the viral RNAs may also comprise the IS sequence, or its complement, described above.

One set of primers for amplification of SARS sequences, particularly by RT-PCR, uses SEQ ID NOs 6562, 6563, 6564 and 6565. Of these, 6562 & 6564 are sense primers and 6563 & 6565 are antisense primers. Primers SEQ ID NOS: 6562 & 6565 may be used in a first amplification, with a second nested amplification being performed using primers SEQ ID NOS: 6563 & 6564. In some embodiments of the invention, these four primers are excluded.

One kit for amplification and detection of SARS sequences, particularly by RT-PCR, uses SEQ ID NOs 6567 & 6568 as primers, and SEQ ID NO 6566 as a probe (typically labeled e.g. with TAMRA and/or FAM) for the amplified sequence. In some embodiments of the invention, these primers and probe are excluded.

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One kit for amplification and detection of SARS sequences, particularly by RT-PCR, uses SEQ ID NOs 7395 & 6568 as primers, and SEQ ID NO 6566 as a probe (typically labeled e.g. with TAMRA and/or FAM) for the amplified sequence. In some embodiments of the invention, these primers and probe are excluded.

One kit for amplification of SARS sequences, particularly the nucleocapsid gene, uses SEQ ID NOs 6560 & 6561 as primers. In some embodiments of the invention, these primers are excluded.

One kit for amplification of SARS sequences uses SEQ ID NOs 6496, 6497, 6562, 6563, 6564 & 6565 as primers. In some embodiments of the invention, these primers are excluded.

One kit for amplification of SARS sequences uses SEQ ID NOs 6562, 6563, 6564 & 6565 as primers. In some embodiments of the invention, these primers are excluded.

One kit for amplification of SARS sequences uses SEQ ID NOs 6500, 6501, 6502 & 6503 as primers. In some embodiments of the invention, these primers are excluded.

One kit for amplification of SARS sequences uses SEQ ID NOs 6496, 6497, 6500, 6501, 6502, 6503, 6562, 6563, 6564 & 6565 as primers. In some embodiments of the invention, these primers are excluded.

One kit for amplification and detection of SARS sequences, particularly by realtime (e.g. TaqManTM) PCR, uses SEQ ID NOs 6567 & 6568 as primers, and SEQ ID NO 6566 as a probe (typically labeled e.g. with TAMRA and/or FAM) for the amplified sequence. In some embodiments of the invention, these primers and probe are excluded.

One kit for amplification and detection of SARS sequences, particularly by realtime (e.g. TaqManTM) PCR, uses SEQ ID NOs 7395 & 6568 as primers, and SEQ ID NO 6566 as a probe (typically labeled e.g. with TAMRA and/or FAM) for the amplified sequence. In some embodiments of the invention, these primers and probe are excluded.

One kit for amplification and detection of SARS sequences uses SEQ ID NOs 6562, 6565 and 6568 as primers, and SEQ ID NOs 7396 and 7397 as probes (typically labeled e.g. with TAMRA and/or FAM) for the amplified sequence. In some embodiments of the invention, these primers and probe are excluded.

One kit for amplification and detection of SARS sequences uses an oligonucleotide comprising SEQ ID NO: 9780 as a forward primer, an oligonucleotide comprising SEQ ID NO: 9781 as a reverse primer, and an oligonucleotide comprising SEQ ID NO: 9782 as a probe.

Preferred sequences for use with RT-PCR and LightCycler analysis include SEQ ID NOs 6562, 6568, 6565, 7396 & 7397. In some embodiments of the invention, these primers and probe are excluded.

The preparation of the oligomers is by means known in the art, including, for example, by methods which include excision, transcription, or chemical synthesis. The target sequences

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and/or regions of the genome which are selected to which the targeting polynucleotides of the oligomers are complementary depend upon the purpose. For example, if the goal is to screen for the presence of SARSV in biological samples (e.g. blood, respiratory material, liver, lung), the preferred oligomers would be used as probes and/or primers, and would hybridize to conserved regions of the SARSV genome. Some of the conserved regions of the SARSV genome to which the oligomers may bind are described herein, for example, 5 UTR and 3 UTR.

In the basic nucleic acid hybridization assay, single-stranded analyte nucleic acid (either DNA or RNA) is hybridized to a nucleic acid probe, and resulting duplexes are detected. The probes for SARSV polynucleotides (natural or derived) are a length which allows the detection of unique viral sequences by hybridization. While 6-8 nucleotides may be a workable length, sequences of 10-12 nucleotides are preferred, and about 13, 14, 15, 16, 17, 18, 19, 20, or 21 or more nucleotides or more appears optimal. Preferably, these sequences will derive from regions which lack heterogeneity. These probes can be prepared using routine methods, including automated oligonucleotide synthetic methods. Among useful probes, for example, are those derived from less conserved regions of the SARSV genome. Regions of the genome that are typically less conserved can be readily ascertained from the sequence alignments provided herein, as well as by any other well known techniques. A complement to any unique portion of the SARSV genome will be satisfactory. For use as probes, complete complementarity is desirable, though it may be unnecessary as the length of the fragment is increased.

For use of such probes as agents to detect the presence of SARSV polynucleotides (for example in screening for contaminated blood or for diagnosing infected individuals), the biological sample to be analyzed, such as, without limitation, blood, serum, lung, liver, mucous, kidney, saliva, or sputum, may be treated, if desired, to extract the nucleic acids contained therein. The resulting nucleic acid from the sample may be subjected to gel electrophoresis or other size separation techniques; alternatively, the nucleic acid sample may be dot blotted without size separation. In order to form hybrid duplexes with the targeting sequence of the probe, the targeted region of the analyte nucleic acid must be in single stranded form. Where the sequence is naturally present in single stranded form, denaturation will not be required. However, where the sequence is present in double stranded form, the sequence will be denatured. Denaturation can be carried out by various techniques known in the art. Subsequent to denaturation, the analyte nucleic acid and probe are incubated under conditions which promote stable hybrid formation of the target sequence in the probe with the putative targeted sequence in the analyte, and the resulting duplexes containing the probe(s) are detected.

Detection of the resulting duplex, if any, is usually accomplished by the use of labeled probes; alternatively, the probe may be unlabeled, but may be detectable by specific binding with a ligand which is labeled, either directly or indirectly. Suitable labels, and methods for labeling

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probes and ligands are known in the art, and include, for example, radioactive labels which may be incorporated by known methods (e.g., nick translation or kinasing), biotin, fluorescent groups, chemiluminescent groups (e.g., dioxetanes, particularly triggered dioxetanes), enzymes, antibodies, and the like.

The region of the probes which are used to bind to the analyte can be made completely complementary to the SARSV genome. Therefore, usually high stringency conditions are desirable in order to prevent false positives. However, conditions of high stringency should only be used if the probes are complementary to regions of the viral genome which lack heterogeneity. The stringency of hybridization is determined by a number of factors during hybridization and during the washing procedure, including temperature, ionic strength, length of time, and concentration of formamide. These factors are outlined in, for example, Maniatis T. (1982).

Variations of this basic scheme which are known in the art, including those which facilitate separation of the duplexes to be detected from extraneous materials and/or which amplify the signal from the labeled moiety, may also be used. A number of these variations are reviewed in, for example: Matthews & Kricka (1988), Analytical Biochemistry 169:1; Landegren et al. (1988), Science 242:229; and Mittlin (1989), Clinical Chem. 35:1819. These and the following publications describing assay formats are hereby incorporated by reference herein. Probes suitable for detecting SARSV in these assays are comprised of sequences which hybridize with target SARSV polynucleotide sequences to form duplexes with the analyte strand, wherein the duplexes are of sufficient stability for detection in the specified assay system.

A suitable variation is, for example, one which is described in US Pat. No. 4,868,105, issued Sep. 9, 1989, and in EPO Publication No. 225,807 (published Jun. 16, 1987). These publications describe a solution phase nucleic acid hybridization assay in which the analyte nucleic acid is hybridized to a labeling probe set and to a capturing probe set. The probe-analyte complex is coupled by hybridization with a solid-supported capture probe that is complementary to the capture probe set. This permits the analyte nucleic acid to be removed from solution as a solid phase complex. Having the analyte in the form of a solid phase complex facilitates subsequent separation steps in the assay. The labeling probe set is complementary to a labeled probe that is bound through hybridization to the solid phase/analyte complex.

The polymerase chain reaction (PCR) is a technique for amplifying a desired nucleic acid sequence (target) contained in a nucleic acid or mixture thereof. In PCR, a pair of primers are employed in excess to hybridize to the complementary strands of the target nucleic acid. The primers are each extended by a polymerase using the target nucleic acid as a template. The extension products become target sequences themselves, following dissociation from the original target strand. New primers then are hybridized and extended by a polymerase, and the cycle is

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repeated to geometrically increase the number of target sequence molecules. PCR is disclosed in US Pat. Nos. 4,683,195 and 4,683,202, which are incorporated herein by reference.

The Ligase Chain Reaction (LCR) is an alternate method for nucleic acid amplification. In LCR, probe pairs are used which include two primary (first and second) and two secondary (third and fourth) probes, all of which are employed in molar excess to target. The first probe hybridizes to a first segment of the target strand, and the second probe hybridizes to a second segment of the target strand, the first and second segments being contiguous so that the primary probes abut one another in 5' phosphate-3' hydroxyl relationship, and so that a ligase can covalently fuse or ligate the two probes into a fused product. In addition, a third (secondary) probe can hybridize to a portion of the first probe and a fourth (secondary) probe can hybridize to a portion of the second probe in a similar abutting fashion. Of course, if the target is initially double stranded, the secondary probes also will hybridize to the target complement in the first instance. Once the ligated strand of primary probes is separated from the target strand, it will hybridize with the third and fourth probes which can be ligated to form a complementary, secondary ligated product. It is important to realize that the ligated products are functionally equivalent to either the target or its complement. By repeated cycles of hybridization and ligation, amplification of the target sequence is achieved. This technique is described more completely in EP-A-320 308 to K. Backman published Jun. 16, 1989 and EP-A-0439182 to K. Backman et al., published Jul. 31, 1991, both of which are incorporated herein by reference.

For amplification of mRNAs, it is within the scope of the present invention to reverse transcribe mRNA into cDNA followed by polymerase chain reaction (RT-PCR); or, to use a single enzyme for both steps as described in US Pat. No. 5,322,770, which is incorporated herein by reference; or reverse transcribe mRNA into cDNA followed by asymmetric gap ligase chain reaction (RT-AGLCR) as described by R. L. Marshall et al., PCR Methods and Applications 4:80-84 (1994), which also is incorporated herein by reference.

TMA is described in detail in, e.g., US Patent No. 5,399,491, the disclosure of which is incorporated herein by reference in its entirety. In one example of a typical assay, an isolated nucleic acid sample, suspected of containing a SARSV target sequence, is mixed with a buffer concentrate containing the buffer, salts, magnesium, nucleotide triphosphates, primers, dithiothreitol, and spermidine. The reaction is optionally incubated at about 100°C for approximately two minutes to denature any secondary structure. After cooling to room temperature, reverse transcriptase, RNA polymerase, and RNAse H are added and the mixture is incubated for two to four hours at 37°C. The reaction can then be assayed by denaturing the product, adding a probe solution, incubating 20 minutes at 60°C, adding a solution to selectively hydrolyze the unhybridized probe, incubating the reaction six minutes at 60°C, and measuring the remaining chemiluminescence in a luminometer.

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Generally, TMA includes the following steps: (a) isolating nucleic acid, including RNA, from the biological sample of interest suspected of being infected with SARSV; and (b) combining into a reaction mixture (i) the isolated nucleic acid, (ii) first and second oligonucleotide primers, the first primer having a complexing sequence sufficiently complementary to the 3' terminal portion of an RNA target sequence, if present (for example the (+) strand), to complex therewith, and the second primer having a complexing sequence sufficiently complementary to the 3' terminal portion of the target sequence of its complement (for example, the (-) strand) to complex therewith, wherein the first oligonucleotide further comprises a sequence 5' to the complexing sequence which includes a promoter, (iii) a reverse transcriptase or RNA and DNA dependent DNA polymerases, (iv) an enzyme activity which selectively degrades the RNA strand of an RNA-DNA complex (such as an RNAse H) and (v) an RNA polymerase which recognizes the promoter.

The components of the reaction mixture may be combined stepwise or at once. The reaction mixture is incubated under conditions whereby an oligonucleotide/target sequence is formed, including DNA priming and nucleic acid synthesizing conditions (including ribonucleotide triphosphates and deoxyribonucleotide triphosphates) for a period of time sufficient to provide multiple copies of the target sequence. The reaction advantageously takes place under conditions suitable for maintaining the stability of reaction components such as the component enzymes and without requiring modification or manipulation of reaction conditions during the course of the amplification reaction. Accordingly, the reaction may take place under conditions that are substantially isothermal and include substantially constant ionic strength and pH. The reaction conveniently does not require a denaturation step to separate the RNA-DNA complex produced by the first DNA extension reaction.

Suitable DNA polymerases include reverse transcriptases, such as avian myeloblastosis virus (AMV) reverse transcriptase (available from, e.g., Seikagaku America, Inc.) and Moloney murine leukemia virus (MMLV) reverse transcriptase (available from, e.g., Bethesda Research Laboratories).

Promoters or promoter sequences suitable for incorporation in the primers are nucleic acid sequences (either naturally occurring, produced synthetically or a product of a restriction digest) that are specifically recognized by an RNA polymerase that recognizes and binds to that sequence and initiates the process of transcription whereby RNA transcripts are produced. The sequence may optionally include nucleotide bases extending beyond the actual recognition site for the RNA polymerase which may impart added stability or susceptibility to degradation processes or increased transcription efficiency. Examples of useful promoters include those which are recognized by certain bacteriophage polymerases such as those from bacteriophage

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T3, T7 or SP6, or a promoter from *E. coli*. These RNA polymerases are readily available from commercial sources, such as New England Biolabs and Epicentre.

Some of the reverse transcriptases suitable for use in the methods herein have an RNAse H activity, such as AMV reverse transcriptase. It may, however, be preferable to add exogenous RNAse H, such as E. coli RNAse H, even when AMV reverse transcriptase is used. RNAse H is readily available from, e.g., Bethesda Research Laboratories.

The RNA transcripts produced by these methods may serve as templates to produce additional copies of the target sequence through the above-described mechanisms. The system is autocatalytic and amplification occurs autocatalytically without the need for repeatedly modifying or changing reaction conditions such as temperature, pH, ionic strength or the like.

Detection may be done using a wide variety of methods, including direct sequencing, hybridization with sequence-specific oligomers, gel electrophoresis and mass spectrometry. these methods can use heterogeneous or homogeneous formats, isotopic or nonisotopic labels, as well as no labels at all.

Suitable labeling moieties for attachment to primers and/or to probes used in methods of

the invention include, but are not limited to: 5-FAM (also called 5-carboxyfluorescein; also called Spiro(isobenzofuran-1(3H), 9'-(9H)xanthene)-5-carboxylic acid,3',6'-dihydroxy-3-oxo-6carboxyfluorescein); 5-Hexachloro-Fluorescein ([4.7.2'.4'.5'.7'-hexachloro-(3'.6'dipivaloylfluoresceinyl)-6-carboxylic acid]); 6-Hexachloro-Fluorescein ([4,7,2',4',5',7'-20 hexachloro-(3',6'-dipivaloylfluoresceinyl)-5-carboxylic acid]); 5-Tetrachloro-Fluorescein ([4,7,2',7'-tetrachloro-(3',6'-dipivaloylfluoresceinyl)-5- carboxylic acid]); 6-Tetrachloro-Fluorescein ([4,7,2',7'-tetrachloro-(3',6'-dipivaloylfluoresceinyl)-6- carboxylic acid]); tetramethylrhodamines (TAMRA), including (i) 5-TAMRA (5-carboxytetramethylrhodamine; Xanthylium, 9-(2,4-dicarboxyphenyl)-3,6- bis(dimethylamino) and (ii) 6-TAMRA (6carboxytetramethylrhodamine; Xanthylium, 9-(2,5-dicarboxyphenyl)-3,6- bis(dimethylamino); 25 EDANS (5-((2-aminoethyl)amino)naphthalene- 1-sulfonic acid); 1,5-IAEDANS (5-(((2-aminoethyl)amino)naphthalene- 1-sulfonic acid); iodoacetyl)amino)ethyl) amino)naphthalene-1-sulfonic acid); DABCYL (4-((4-(dimethylamino)phenyl) azo)benzoic acid); Cv5 (Indodicarbocyanine-5); Cv3 (Indodicarbocyanine-3); and BODIPYTM FL (4,4-difluoro-5,7-dimethyl-4-bora-3a,4a-diaza-s-

Nucleic acids of the invention may be used in solution or may be bound to a solid matrix or support e.g. in the format of a DNA array,

indacene-3-propionic acid). Labelling of probes with both FAM (e.g. at 5') and TAMRA (e.g. at

As is readily apparent, design of the assays described herein are subject to a great deal of variation, and many formats are known in the art. The above descriptions are merely provided as

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3') is preferred.

guidance and one of skill in the art can readily modify the described protocols, using techniques well known in the art.

One 302nt amplicon of the SARS virus is known as "BNI-1" (SEQ ID NO: 9927). It was sequenced at the Bernhard Nocht Institute, Hamburg, Germany. In April 2003 the BNI-1 sequence was published on the WHO website (http://www.who.int/csr/sars/primers/en/) and in Dorsten et al., "Identification of a Novel Coronavirus in Patients with Severe Acute Respiratory Syndrome", New England Journal of Medicine, published online at http://www.nejm.org. Both references are incorporated herein by reference in their entirety. Some embodiments of the invention do not encompass a nucleic acid consisting of SEQ ID NO: 9927. Some other embodiments of the invention do not encompass a nucleic acid comprising SEQ ID NO: 9927. Some embodiments of the invention do not encompass a polypeptide consisting of any one of SEQ ID NO. Sep 28 to 9959. Some other embodiments of the invention do not encompass a nucleic acid comprising any one of SEQ ID NO. Sep 28 to 9959. Some embodiments of the invention are not subject to these exclusions.

5 Immunoassays

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The present invention utilizes various immunoassay techniques for identifying individuals exposed to SARSV and/or biological samples containing SARSV antigens or antibodies to SARSV.

Immunoassay Formats

The SARSV antigens may be employed in virtually any assay format that employs a known antigen to detect antibodies. A common feature of all of these assays is that the antigen is contacted with biological sample suspected of containing SARSV antibodies under conditions that permit the antigen to bind to any such antibody present in the component. Such conditions will typically be physiologic temperature, pH and ionic strength using an excess of antigen. The incubation of the antigen with the specimen is followed by detection of immune complexes comprised of the antigen. Alternatively, anti-SARSV antibodies may be employed to detect the presence of SARSV antigens in a biological sample. Combination antigen/antibody assays are also contemplated; for example, as described for HCV detection in US patent 6,630,298.

Design of the immunoassays is subject to a great deal of variation, and many formats are known in the art. Protocols may, for example, use solid supports, or immunoprecipitation. Most assays involve the use of labeled antibody or polypeptide; the labels may be, for example, enzymatic, fluorescent, chemiluminescent, radioactive, or dye molecules. Assays which amplify the signals from the immune complex are also known; examples of which are assays which utilize biotin and avidin, and enzyme-labeled and mediated immunoassays, such as ELISA assays.

The immunoassay may be, without limitation, in a heterogeneous or in a homogeneous format, and of a standard or competitive type. In a heterogeneous format, the polypeptide is typically bound to a solid matrix or support to facilitate separation of the sample from the polypeptide after incubation. Examples of solid supports that can be used are nitrocellulose (e.g., in membrane or microtiter well form), polyvinyl chloride (e.g., in sheets or microtiter wells), polystyrene latex (e.g., in beads or microtiter plates, polyvinylidine fluoride, diazotized paper, nylon membranes, microchips, high or low density biochips, recombinant immunoassays (RIBA), microfluidity devices, micromagnetic beads, activated beads, and Protein A beads. For example, Dynatech Immunlon or Immunlon 2 microtiter plates or 0.25 inch polystyrene beads (Precision Plastic Ball) can be used in the heterogeneous format. The solid support containing the antigenic polypeptides is typically washed after separating it from the test sample, and prior to detection of bound antibodies. Both standard and competitive formats are known in the art.

In a homogenous format, the test sample is incubated with the combination of antigens in solution. For example, it may be under conditions that will precipitate any antigen-antibody complexes which are formed. Both standard and competitive formats for these assays are known in the art.

In a standard format, the amount of SARSV antibodies in the antibody-antigen complexes is directly monitored. This may be accomplished by determining whether labeled anti-xenogeneic (e.g., anti-human) antibodies which recognize an epitope on anti-SARSV antibodies will bind due to complex formation. In a competitive format, the amount of SARSV antibodies in the sample is deduced by monitoring the competitive effect on the binding of a known amount of labeled antibody (or other competing ligand) in the complex.

Complexes formed comprising anti-SARSV antibody (or in the case of competitive assays, the amount of competing antibody) are detected by any of a number of known techniques, depending on the format. For example, unlabeled SARSV antibodies in the complex may be detected using a conjugate of antixenogeneic Ig complexed with a label, (e.g., an enzyme label).

In an immunoprecipitation or agglutination assay format the reaction between the SARSV antigens and the antibody forms a network that precipitates from the solution or suspension and forms a visible layer or film of precipitate. If no anti-SARSV antibody is present in the test specimen, no visible precipitate is formed.

There are at least three specific types of particle agglutination (PA) assays. These assays are used for the detection of antibodies to various antigens when coated to a support. One type of this assay is the hemagglutination assay using red blood cells (RBCs) that are sensitized by passively adsorbing antigen (or antibody) to the RBC. The addition of specific antigen antibodies present in the body component, if any, causes the RBCs coated with the purified antigen to agglutinate.

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To eliminate potential non-specific reactions in the hemagglutination assay, two artificial carriers may be used instead of RBC in the PA. The most common of these are latex particles. However, gelatin particles may also be used. The assays utilizing either of these carriers are based on passive agglutination of the particles coated with purified antigens.

The SARSV antigens will typically be packaged in the form of a kit for use in these immunoassays. The kit will normally contain in separate containers the native SARSV antigen, control antibody formulations (positive and/or negative), labeled antibody when the assay format requires same and signal generating reagents (e.g., enzyme substrate) if the label does not generate a signal directly. The native SARSV antigen may be already bound to a solid matrix or separate with reagents for binding it to the matrix. Instructions (e.g., written, tape, CD-ROM, etc.) for carrying out the assay usually will be included in the kit.

Immunoassays that utilize the native SARSV antigen are additionally useful in screening blood for the preparation of a supply from which potentially infective SARSV is lacking. The method for the preparation of the blood supply comprises the following steps. Reacting a body component, preferably blood or a blood component, from the individual donating blood with native SARSV antigen to allow an immunological reaction between SARSV antibodies, if any, and the SARSV antigen. Detecting whether anti-SARSV antibody—SARSV antigen complexes are formed as a result of the reacting. Blood contributed to the blood supply is from donors that do not exhibit antibodies to the native SARSV antigens.

Production of Antibodies

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As explained above, the assay may utilize various antibodies which may be bound to a solid support, and that detect antigen or antigen/antibody complexes formed when SARSV infection is present in the sample. These antibodies may be polycional or monoclonal antibody preparations, monospecific antisera, human antibodies, or may be hybrid or chimeric antibodies, such as humanized antibodies, altered antibodies, F(ab')₂ fragments, F(ab) fragments, Fv fragments, single-domain antibodies, dimeric or trimeric antibody fragment constructs, minibodies, or functional fragments thereof which bind to the antigen in question.

Antibodies are produced using techniques well known to those of skill in the art and disclosed in, for example, US Pat. Nos. 4,011,308; 4,722,890; 4,016,043; 3,876,504; 3,770,380; and 4,372,745. For example, polyclonal antibodies are generated by immunizing a suitable animal, such as a mouse, rat, rabbit, sheep or goat, with an antigen of interest. In order to enhance immunogenicity, the antigen can be linked to a carrier prior to immunization. Such carriers are well known to those of ordinary skill in the art. Immunization is generally performed by mixing or emulsifying the antigen in saline, preferably in an adjuvant such as Freund's complete adjuvant, and injecting the mixture or emulsion parenterally (generally subcutaneously or intramuscularly). The animal is generally boosted 2-6 weeks later with one or more injections

of the antigen in saline, preferably using Freund's incomplete adjuvant. Antibodies may also be generated by *in vitro* immunization, using methods known in the art. Polyclonal antiserum is then obtained from the immunized animal.

Monoclonal antibodies are generally prepared using the method of Kohler & Milstein (1975) *Nature* 256:495-497, or a modification thereof, as described above.

As explained above, antibody fragments which retain the ability to recognize the antigen of interest, will also find use in the subject immunoassays. A number of antibody fragments are known in the art which comprise antigen-binding sites capable of exhibiting immunological binding properties of an intact antibody molecule. For example, functional antibody fragments can be produced by cleaving a constant region, not responsible for antigen binding, from the antibody molecule, using e.g., pepsin, to produce F(ab')₂ fragments. These fragments will contain two antigen binding sites, but lack a portion of the constant region from each of the heavy chains. Similarly, if desired, Fab fragments, comprising a single antigen binding site, can be produced, e.g., by digestion of polyclonal or monoclonal antibodies with papain. Functional fragments, including only the variable regions of the heavy and light chains, can also be produced, using standard techniques such as recombinant production or preferential proteolytic cleavage of immunoglobulin molecules. These fragments are known as Fv. See, e.g., Inbar et al. (1972) Proc. Nat. Acad. Sci. USA 69:2659-2662; Hochman et al. (1976) Biochem 15:2706-2710; and Ehrlich et al. (1980) Biochem 19:4091-4096.

A single-chain Fv ("sFv" or "scFv") polypeptide is a covalently linked V_H-V_L heterodimer which is expressed from a gene fusion including V_H- and V_L-encoding genes linked by a peptideencoding linker. Huston et al. (1988) Proc. Nat. Acad. Sci. USA 85:5879-5883. A number of methods have been described to discern and develop chemical structures (linkers) for converting the naturally aggregated, but chemically separated, light and heavy polypeptide chains from an antibody V region into an sFv molecule which will fold into a three dimensional structure substantially similar to the structure of an antigen-binding site. See, e.g., US Pat. Nos. 5,091,513, 5,132,405 and 4,946,778. The sFv molecules may be produced using methods described in the art. See, e.g., Huston et al. (1988) Proc. Nat. Acad. Sci. USA 85:5879-5883; US Pat. Nos. 5,091,513, 5,132,405 and 4,946,778. Design criteria include determining the appropriate length to span the distance between the C-terminus of one chain and the N-terminus of the other, wherein the linker is generally formed from small hydrophilic amino acid residues that do not tend to coil or form secondary structures. Such methods have been described in the art. See, e.g., US Pat. Nos. 5,091,513, 5,132,405 and 4,946,778. Suitable linkers generally comprise polypeptide chains of alternating sets of glycine and serine residues, and may include glutamic acid and lysine residues inserted to enhance solubility.

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"Mini-antibodies" or "minibodies" will also find use with the present invention. Minibodies are sFv polypeptide chains which include oligomerization domains at their C-termini, separated from the sFv by a hinge region. Pack et al. (1992) Biochem 31:1579-1584. The oligomerization domain comprises self-associating a-helices, e.g., leucine zippers, that can be further stabilized by additional disulfide bonds. The oligomerization domain is designed to be compatible with vectorial folding across a membrane, a process thought to facilitate in vivo folding of the polypeptide into a functional binding protein. Generally, minibodies are produced using recombinant methods well known in the art. See, e.g., Pack et al. (1992) Biochem 31:1579-1584; Cumber et al. (1992) J. Immunology 149B: 120-126.

10 Production of SARS Antigens

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The SARSV antigens used in the present invention are generally produced recombinantly. Thus, polynucleotides encoding SARSV antigens for use with the present invention can be made using standard techniques of molecular biology. For example, polynucleotide sequences coding for the above-described molecules can be obtained using recombinant methods, such as by screening cDNA and genomic libraries from cells expressing the gene, or by deriving the gene from a vector known to include the same. Furthermore, the desired gene can be isolated directly from viral nucleic acid molecules, using techniques described in the art, such as those described for HCV in Houghton et al., US Pat. No. 5,350,671. The gene encoding the antigen of interest can also be produced synthetically, rather than cloned. The molecules can be designed with appropriate codons for the particular sequence (preferably optimum codons for the expression host of choice). The complete sequence is then assembled from overlapping oligonucleotides prepared by standard methods and assembled into a complete coding sequence. See, e.g., Edge (1981) Nature 292:756; Nambair et al. (1984) Science 223:1299; and Jay et al. (1984) J. Biol. Chem. 259:6311.

Thus, particular nucleotide sequences can be obtained from vectors harboring the desired sequences or synthesized completely or in part using various oligonucleotide synthesis techniques known in the art, such as site-directed mutagenesis and polymerase chain reaction (PCR) techniques where appropriate. See, e.g., Sambrook, supra. In particular, one method of obtaining nucleotide sequences encoding the desired sequences is by annealing complementary sets of overlapping synthetic oligonucleotides produced in a conventional, automated polynucleotide synthesizer, followed by ligation with an appropriate DNA ligase and amplification of the ligated nucleotide sequence via PCR. See, e.g., Jayaraman et al. (1991) Proc. Natl. Acad. Sci. USA 88:4084-4088. Additionally, oligonucleotide directed synthesis (Jones et al. (1986) Nature 54:75-82), oligonucleotide directed mutagenesis of pre-existing nucleotide regions (Riechmann et al. (1988) Nature 332:323-327 and Verhoeyen et al. (1988)

Science 239:1534-1536), and enzymatic filling-in of gapped oligonucleotides using T4 DNA polymerase (Queen et al. (1989) Proc. Natl. Acad. Sci. USA 86:10029-10033) can be used under the invention to provide molecules having altered or enhanced antigen-binding capabilities, and/or reduced immunogenicity.

Once coding sequences have been prepared or isolated, such sequences can be cloned into any suitable vector or replicon. Numerous cloning vectors are known to those of skill in the art, and the selection of an appropriate cloning vector is a matter of choice. Suitable vectors include, but are not limited to, plasmids, phages, transposons, cosmids, chromosomes (including artificial chromosomes, such as BACs or YACs) or viruses which are capable of replication when associated with the proper control elements.

The coding sequence is then placed under the control of suitable control elements, depending on the system to be used for expression. Thus, the coding sequence can be placed under the control of a promoter, ribosome binding site (for bacterial expression) and, optionally, an operator, so that the DNA sequence of interest is transcribed into RNA by a suitable transformant. The coding sequence may or may not contain a signal peptide or leader sequence which can later be removed by the host in post-translational processing. See, e.g., US Pat. Nos. 4,431,739; 4,425,437; 4,338,397.

In addition to control sequences, it may be desirable to add regulatory sequences which allow for regulation of the expression of the sequences relative to the growth of the host cell. Regulatory sequences are known to those of skill in the art, and examples include those which cause the expression of a gene to be turned on or off in response to a chemical or physical stimulus, including the presence of a regulatory compound. Other types of regulatory elements may also be present in the vector. For example, enhancer elements may be used herein to increase expression levels of the constructs. Examples include the SV40 early gene enhancer (Dijkema et al. (1985) EMBO J. 4:761), the enhancer/promoter derived from the long terminal repeat (LTR) of the Rous Sarcoma Virus (Gorman et al. (1982) Proc. Natl. Acad. Sci. USA 79:6777) and elements derived from human CMV (Boshart et al. (1985) Cell 41:521), such as elements included in the CMV intron A sequence (US Pat. No. 5,688,688). The expression cassette may further include an origin of replication for autonomous replication in a suitable host cell, one or more selectable markers, one or more restriction sites, a potential for high copy number and a strong promoter.

An expression vector is constructed so that the particular coding sequence is located in the vector with the appropriate regulatory sequences, the positioning and orientation of the coding sequence with respect to the control sequences being such that the coding sequence is transcribed under the "control" of the control sequences (i.e., RNA polymerase which binds to the DNA molecule at the control sequences transcribes the coding sequence). Modification of the

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sequences encoding the molecule of interest may be desirable to achieve this end. For example, in some cases it may be necessary to modify the sequence so that it can be attached to the control sequences in the appropriate orientation; i.e., to maintain the reading frame. The control sequences and other regulatory sequences may be ligated to the coding sequence prior to insertion into a vector. Alternatively, the coding sequence can be cloned directly into an expression vector which already contains the control sequences and an appropriate restriction site

As explained above, it may also be desirable to produce mutants or analogs of the antigen of interest. Methods for doing so are described in, e.g., Dasmahapatra et al., US Pat. No. 5.843,752 and Zhang et al., US Pat. No. 5,990,276. Mutants or analogs of SARSV proteins for use in the subject assays may be prepared by the deletion of a portion of the sequence encoding the polypeptide of interest, by insertion of a sequence, and/or by substitution of one or more nucleotides within the sequence. Techniques for modifying nucleotide sequences, such as sitedirected mutagenesis, and the like, are well known to those skilled in the art. See, e.g., Sambrook et al., supra; Kunkel, T. A. (1985) Proc. Natl. Acad. Sci. USA (1985) 82:448; Geisselsoder et al. (1987) BioTechniques 5:786; Zoller & Smith (1983) Methods Enzymol. 100:468; Dalbie-McFarland et al. (1982) Proc. Natl. Acad. Sci USA 79:6409.

The molecules can be expressed in a wide variety of systems, including insect, mammalian, bacterial, viral and yeast expression systems, all well known in the art.

For example, insect cell expression systems, such as baculovirus systems, are known to those of skill in the art and described in, e.g., Summers & Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987). Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, inter alia, Invitrogen, San Diego Calif. ("MaxBac" kit). Similarly, bacterial and mammalian cell expression systems are well known in the art and described in, e.g., Sambrook et al., supra. Yeast expression systems are also known in the art and described in, e.g., Yeast Genetic Engineering (Barr et al., eds., 1989) Butterworths, London.

A number of appropriate host cells for use with the above systems are also known. For example, mammalian cell lines are known in the art and include immortalized cell lines available from the American Type Culture Collection (ATCC), such as, but not limited to, Chinese) hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (COS), human embryonic kidney cells, human hepatocellular carcinoma cells (e.g., Hep G2), Madin-Darby bovine kidney ("MDBK") cells, as well as others. Similarly, bacterial hosts such as E.coli, Bacillus subtilis, and Streptococcus spp., will find use with the present expression constructs. Yeast hosts useful in the present invention include inter alia, Saccharomyces cerevisiae, Candida albicans, Candida maltosa, Hansenula polymorpha, Kluyveromyces fragilis,

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Kluyveromyces lactis, Pichia guillerimondii, Pichia pastoris, Schizosaccharomyces pombe and Yarrowia lipolytica. Insect cells for use with baculovirus expression vectors include, inter alia, Aedes aegypti, Autographa califormica, Bombyx mori, Drosophila melanogaster, Spodoptera frugiperda, and Trichoplusia ni.

Nucleic acid molecules comprising nucleotide sequences of interest can be stably integrated into a host cell genome or maintained on a stable episomal element in a suitable host cell using various gene delivery techniques well known in the art. See, e.g., US Pat. No. 5,399,346.

Depending on the expression system and host selected, the molecules are produced by growing host cells transformed by an expression vector described above under conditions whereby the protein is expressed. The expressed protein is then isolated from the host cells and purified. If the expression system secretes the protein into growth media, the product can be purified directly from the media. If it is not secreted, it can be isolated from cell lysates. The selection of the appropriate growth conditions and recovery methods are within the skill of the art.

EXAMPLE ...

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For useful expression of SARSV antigens in Saccharomyces cerevisiae and Pichia pastoris, insect cells, and mammalian cells, the following domains are cloned into expression vectors as listed in the Table below. The nt sequence numbers are from the SARSV sequence of SEO ID NO: 1.

- RNA polymerase 1a: SARS nt 250-13398
- RNA polymerase 1b: SARS nt 13399-21470
- ORFns.envelope (homologous to ns2, hemagglutinin-esterase envelope glycoprotein, and spike glycoprotein): SARS nt 21477-25244
- Membrane: SARS nt 27849 28103
 - Nucleocapsid: SARS nt 28105 29373

A combination of PCR and synthetic oligos is used to create the above domains with restriction sites tailored to the following expression vectors:

Restriction ends HindIII/SalI EcoRI/Sal\\\ Xbal/SalI	Vector pBS24.1 pBS24.1 pAO815	Promoter ADH2/GAPDH ADH2/GAPDH/SOD fusion AOXI	Expression host AD3/Saccharomyces AD3/Saccharomyces GS115/Pichia pastoris
EcoRI/BamHI EcoRI/XmaI	pCMVkm2 pCMVIII	CMVp/Enhancer/IntronA CMVp/Enhancer/IntronA	HVK-293/Transient transfection CHO stable cell line
NheI/SaII	pBluBac4.5	Polyhedrin	Cell lines employed by Chiron include: Sf9, Sf21, Tn5

IV. TREATMENT OF SARS INFECTION WITH RNAi

RNA interference or "RNAi" is a term initially coined by Fire and co-workers to describe the observation that double-stranded RNA (dsRNA) can block gene expression when it is introduced into worms (Fire et al., Nature 391, 806-811(1998)). RNAi most likely involves mRNA degradation, resulting in sequence-specific, post-transcriptional gene silencing in many organisms. RNAi is a post-transcriptional process triggered by the introduction of double-stranded RNA which leads to gene silencing in a sequence-specific manner. RNAi has been reported to occur naturally in organisms as diverse as nematodes, trypanosmes, plants and fungi. It most likely serves to protect organisms from viruses, modulate transposon activity and eliminate aberrant transcription products.

The first evidence that dsRNA could achieve efficient gene silencing through RNAi came from studies on the nematode *Caenorhabditis elegans* (Fire et al. (1998) *Nature*, 391:806-811 and US Patent No. 6,506,559). Later studies in the fruit fly *Drosophila melanogaster* demonstrated that RNAi is a two-step mechanism (Elbashir et al. (2001) *Genes Dev.*, 15(2): 188-200). First, long dsRNAs are cleaved by an enzyme known as Dicer in 21-23 nucleotides (nt) fragments, called small interfering RNAs (siRNAs). Then, siRNAs associate with a ribonuclease complex (termed RISC for RNA Induced Silencing Complex) which target this complex to complementary mRNAs. RISC then cleaves the targeted mRNAs opposite the complementary siRNA, which makes the mRNA susceptible to other RNA degradation pathways.

RNAi is the phenomenon where dsRNA corresponding to a targeted DNA or RNA sequence can suppress or silence gene expression. Even though dsRNA can mediate gene-specific interference in mammalian cells in some circumstances (Wianny & Zernicka-Goetz (2000) Nature Cell Biol. 2:70-75; Svoboda et al. (2000) Development 17:4147-4156) the use of RNAi in mammalian somatic cells is often limited due to the dsRNA triggering dsRNA-dependent protein kinase (PKR) which in turn inactivates translation factor eIF2a and causes a generalized suppression of protein synthesis and often times apoptosis (Gil & Esteban (2000) Apoptosis 5:107-114).

Recently, gene-specific suppression using siRNA of approximately 21 or 22 base pairs in length, corresponding to targeted RNA or DNA sequences, were shown to disrupt the expression of these targeted sequences in mammalian cells (Elbashir, S.M., et al., Nature 411: 494-498 (2001)). However, it is not clear that all RNA or DNA sequences of a mammalian cell's genome are susceptible to siRNA. It is also uncertain that every mammalian cell type possesses the necessary machinery for effecting gene-specific suppression using siRNA. Further, siRNA is of limited use for at least two reasons: the transient nature of the suppression effect seen in cells where the siRNA has been administered; and in some instances the necessity for chemical synthesis of siRNAs before their use (Tuschl T., Nature Biotechnol., 20: 446-448 (2002)). Also

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the instability of these short, synthetic RNAs makes it presents problems for any long term use of these siRNAs a pharmaceutical.

To overcome this limitation, the present invention provides a modified siRNA with increased stability against nuclease degradation while still maintaining its ability to inhibit viral replication via RNA interference. Such modification to the ribonucleotides in the siRNAs, adds a chemical group via chemical synthesis or *in vitro* transcription or longer modified RNAs can be prepared by either of these methods and cut into siRNAs using Dicer.

Although other methods for gene-specific suppression have utilized chemically-modified nucleic acids, such as antisense and ribozyme technology, such modification destroys critical enzymatic activities necessary for the function of these technologies. In regard to antisense technology, modification of the ribonucleotides destroys RNaseH activity, whereas such modification abolishes the catalytic activity of ribozymes.

The present invention provides a double-stranded RNA (dsRNA) molecule modified for protection against nuclease degradation with a length from about 10 to about 30 nucleotides which is able to inactivate a virus in a mammalian cell. The invention also provides a method of inactivating a virus by administering modified small interfering RNAs (siRNAs) that are modified so that they are nuclease or RNase resistant and retain the biological activity of being able to inhibit viral replication by targeting a RNA sequence in a virus.

The invention is further directed to a method of making modified siRNAs that target a RNA sequence in a virus comprising preparing a modified-double stranded RNA (dsRNA) fragment containing at least one modified ribonucleotide in at least one strand that spans the genome of the virus; and cleaving the modified-dsRNA fragments with recombinant human Dicer resulting in more than one modified siRNA.

The present invention provides a modified dsRNA molecule of from about 10 to about 30 nucleotides which mediates targeted RNA interference in hepatic or SARS-infected cells.

As used herein RNA interference, or RNAi, is used to mean sequence-specific, or gene specific, suppression of gene expression (protein synthesis), without causing a generalized suppression of protein synthesis in cells harboring the siRNA. The invention is not limited to a particular theory of the mechanism of action of RNAi. For example, RNAi may involve degradation of messenger RNA (mRNA) in an RNA-induced silencing complex (RISC), preventing translation of the transcribed mRNA, or it may involve the methylation of genomic DNA, shunting transcription of the gene. The lack of gene expression caused by RNAi may be transient, lasting a short period of time, or it may be stable, or permanent, lasting an indefinite period of time.

The term RNA is meant as is recognized in the art. Further, as used herein, RNA is used to mean double-stranded RNA (dsRNA) or single-stranded RNA (ssRNA) or a dsRNA with a

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single-stranded overhang. dsRNAs within the meaning of the present invention includes short interfering RNA (siRNA), micro RNA (miRNA) and small hairpin RNA (shRNA), Additionally, RNA is also used to mean messenger RNA (mRNA), transfer RNA (tRNA) or ribosomal RNA (tRNA).

The present invention is directed to small interfering RNA (siRNA) which have been chemically modified to confer increased stability against nuclease degradation yet these siRNAs are still able to bind to target RNAs, that may be present in a cells. In the case where the target RNA is a virus specific RNA, the modified siRNAs are able to bind to the virus specific RNAs and inactivate the virus. A modified siRNA of the present invention comprises a modified ribonucleotide, wherein the siRNA is resistant to enzymatic degradation, such as RNase degradation, and yet retains the ability to inhibit viral replication. The modified siRNA is more specifically modified at the 2' position of the ribose in the siRNA. The modification is at the 2' position of at least one ribonucleotide of said siRNA. Attachment of receptor-binding ligands to siRNA molecules can be used to target the siRNA to a desired cell type. For example, attachment of cholesterol at the 5'-end or 3'-end of the siRNA molecule, to give a cholesteryl siRNA, can enhance targeting to hepatocytes. Other ligands for receptor mediated siRNA targeting to liver include HBV surface antigen, LIDL, and others.

More specifically, the siRNA is modified at at least one pyrimidine, at least one purine or a combination thereof. However, generally all pyrimidines, or all purines or a combination of all pyrimidines and all purines of the siRNA are modified. More preferably, the pyrimidines are modified and these pyrimidines are cytosine, a derivative of cytosine, uracil, a derivative of uracil or a combination thereof. It also is contemplated to modify the selected ribonucleotides in at least one strand of the siRNA or the ribonucleotides in both strands of the siRNA are modified.

The nucleotides containing pyrimidine bases found in RNA (cytidine and uridine) can be chemically modified by adding any molecule that inhibits RNA degradation or breakdown to the 2' position of the ribose molecule. The 2'-modified pyrimidine nucleotide can be formed using a number of different methods. The 2' modification confers increased stability to the siRNA by making the siRNA impervious or resistant to nuclease activity. Thus, the 2' modified siRNA has a longer serum half-life and is resistant to degradation compared to unmodified siRNA. The siRNA also may be modified completely or partially.

Regarding chemical modification of siRNAs, a molecule from the halide chemical group is preferably added to the ribonucleotide of the siRNA. Within the halides, fluorine is the preferred molecule but other chemical molecules, in addition to fluoro-, such as methyl-, methoxyethyl- and propyl-modifications can also we made. But the preferred modications is fluoro-modification, such as a 2'-fluoro-modication or a 2',2'-fluoro-modification. Thus, in a preferred

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embodiment of the invention, the siRNA is modified by adding a fluorine molecule to the 2' carbon of the pyrimidine ribonucleotide. The siRNA may be fluorinated completely or partially. For example, only the cytosine nucleotides need be fluorinated. Alternatively, only the uracil nucleotide need be fluorinated but both uracil and cytosine can be fluorinated. Furthermore, only one strand, either sense or antisense, of the siRNA can be fluorinated. Even partial 2' fluorination the siRNA gives protection against nucleolytic degradation. Furthermore, it is important to note the 2' fluorinated siRNA is not toxic to cells, an unexpected result given that fluorine chemistry usually is toxic to living organisms.

The siRNA of the present invention is designed to interact with a target nucleotide sequence. Most preferably this target nucleotide sequence is a disease producing agent or pathogen of which one wishes to inhibit gene expression. More preferably, this target nucleotide sequence is in a virus genome, and further this virus genome is from a RNA virus or a DNA virus is selected from the group consisting of hepatitis C virus (HCV), hepatitis A virus, hepatitis B virus, hepatitis D virus, hepatitis E virus, Ebola virus, influenza virus, rotavirus, retrovirus, poliovirus, human papilloma virus (HPV), metapneumovirus and coronavirus. The most preferred virus is SARS virus.

Modfied siRNA may be prepared in a number of ways, such as by chemical synthesis, T7 polymerase transcription, or by treating modified long double stranded RNA (dsRNA) prepared by one of the two previous methods with Dicer enzyme. Dicer enzyme can be used to cleave dsRNA that is about 500 base pairs to about 1000 base pairs in size, to created mixed populations of dsRNA from about 21 to about 23 base pairs in length. Furthermore, an unexpected result of using the Dicer enzyme method is that Dicer enzyme will cleave modified strands of dsRNA, such as 2' fluorinated modified dsRNA. Before development of this method, it was previously thought that Dicer would not be able to cleave modified siRNA. The Dicer method can be carried out using the Dicer siRNA Generation Kit available from Gene Therapy Systems, San Diego, CA.

As used herein, small interfering RNA (siRNA) is defined as double- or single-stranded RNA of from about 10 to about 30 nucleotides in length, more preferably 12-28 nucleotides, more preferably 15-25 nucleotides, even more preferably 19-23 nucleotides and most preferably 21-23 nucleotides. The length of a siRNA as used herein, is determined by the length of one of the strands of the RNA. For example, a siRNA that is described as 21 nucleotides long (a 21-mer) may comprise two opposite strands of RNA which anneal together for 19 contiguous base pairings. The two remaining nucleotides on one end of the molecule would not anneal to the opposite strand, thus creating an "overhang". The overhang can be at the 5' or the 3' end of the dsRNA. Preferably, the overhang is at the 3' end of the RNA strand. The length of a double-stranded RNA where the two opposite strands are not the same length will be designated by the

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longer of the two strands. For example, a dsRNA comprising one strand which is 21 nucleotides long and anneals to an opposite strand that is 20 nucleotides long, will be considered, as used herein, a 21-mer.

Preferably, the siRNA of the present invention will comprise a 3' overhang of about 2 to 4 bases. More preferably, the 3' overhang is 2 nucleotides long. Even more preferably, the 2 nucleotides comprising the 3' overhang are uridine (U).

In one embodiment, the invention provides an RNA molecule comprising a nucleotide sequence at least 80% identical to the nucleotide sequence of the target agent or virus. Preferably, the RNA molecule of the present invention is at least 90%, 95%, 96%, 97%, 98%, 99% or 100% identical to the nucleotide sequence of the target agent or virus.

As a practical matter, whether any particular nucleic acid molecule is at least 90%, 95%, 96%, 97% 98%, 99% or 100% identical to the nucleotide sequence of the target agent or virus can be determined conventionally using known computer programs such as the *Bestfit* program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, Wis. 53711). *Bestfit* uses the local homology algorithm of Smith & Waterman (*Advances in Applied Mathematics* 2:482-489 (1981)) to find the best segment of homology between two sequences. When using *Bestfit* or any other sequence alignment program to determine whether a particular sequence is, for instance, 95% identical to a reference sequence according to the present invention, the parameters are set, of course, such that the percentage of identity is calculated over the full length of the reference nucleotide sequence and that gaps in homology of up to 5% of the total number of nucleotides in the reference sequence are allowed.

The present invention provides a method of inactivating a target agent or preferably a virus in a patient comprising administering to the patient a modified siRNA in an effective amount to inactivate the targeted agent or virus. RNA interference towards a targeted DNA segment in a cell can be achieved by administering a dsRNA molecule or siRNA to the cells, wherein the nucleotide sequence of the dsRNA molecule corresponds to the nucleotide sequence of the targeted DNA segment. Preferably, the RNA molecule used to induce targeted RNAi is siRNA.

Gene suppression, targeted suppression, sequence-specific suppression, targeted RNAi or sequence-specific RNAi are used interchangeably herein. Furthermore, sequence-specific suppression, as used herein, is determined by separately assaying the levels of the protein targeted for suppression in cells containing the siRNA (experimental cells) and in cells not containing the identical siRNA (control cells), and comparing the two values. Furthermore, the experimental and control cells must be derived from the same source and same animal. For example, the control and experimental cells can be, but are not limited to, normal human hepatic cells as cell culture *in vitro*, or they can derived from a hepatocellular carcinoma. Further, the

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control and experimental cells used in determining the level or quantity of gene suppression must be assayed under similar, if not identical, conditions.

As used herein the phrase "targeted DNA segment" is used to mean a DNA sequence encoding, in whole or in part, an mRNA for a targeted protein, including introns or exons, where suppression is desired. DNA segment can also mean a DNA sequence that normally regulates expression of the targeted protein, including but not limited to the promoter of the targeted protein. Furthermore, the DNA segment may or may not be a part of the cell's genome or it may be extrachromosomal, such as plasmid DNA.

The present invention is further directed to inactivating a virus in a patient comprising administering to a patient a modified siRNA in an effective amount to inactivate the virus. The siRNA is preferably about 10 to about 30 nucleotides in length, more preferably 12-28 nucleotides, more preferably 15-25 nucleotides, even more preferably 19-23 nucleotides and most preferably 21-23 nucleotides. The method preferably utilizes a 2' modified siRNA that is modified at the 2' position of at least one ribonucleotide of said siRNA. The method utilizes a siRNA that is modified with chemical groups selected from the group consisting of fluoro-, methyl-, methoxyethyl- and propyl-modification. The fluoro-modification is preferred and either a 2'-fluoro-modication or a 2',2'-fluoro-modification is useful in the present invention and preferred.

The modification may be at the pyrimidines, the purines or a combination thereof of the siRNA are modified. More preferably the pyrimidines are modified, such as cytosine, a derivative of cytosine, uracil, a derivative of uracil or a combination thereof. In one embodiment, at least one strand of the siRNA contains at least one modified nucleotide and in an alternate embodiment, oth strands of the siRNA contains at least one modified nucleotide.

The method is intended to target disease causing agents or pathogens, an more particularly viruses, which can be either a RNA virus or a DNA virus, which are selected from the group consisting of hepatitis C virus (HCV), hepatitis A virus, hepatitis B virus, hepatitis D virus, hepatitis E virus, Ebola virus, influenza virus, rotavirus, recovirus, retrovirus, poliovirus, human papilloma virus (HPV), metapneumovirus and coronavirus. More preferably the target virus is a SARS virus. The present method utilizes a siRNA prepared by (a) identifying a target nucleotide sequence in a virus genome, preferably SARS virus, for designing a small interfering RNA (siRNA); and (b) producing a siRNA that has been modified to contain at least one modified nucleotide. More preferably, the siRNA comprises a dsRNA molecule with a first strand ribonucleotide sequence corresponding to a nucleotide sequence corresponding to a target nucleotide sequence in said virus and a second strand comprising a ribonucleotide sequence complementary to said target nucleotide sequence, wherein said first and second strands are separate complementary strands that hybridize to each other to form said dsRNA molecule, and

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further wherein the first strand ribonucleotide sequence, the second strand ribonucleotide sequence or both the first and second strand ribonucletide sequences comprise at least one modified nucleotide. In this method, the target nucleotide sequence comprises a conserved nucleotide sequence necessary for SARS virus replication, and the conserved nucleotide sequence is selected from the group consisting of SEQ ID NO: 7292, SEQ ID NO: 7293, SEQ ID NO: 7294, SEQ ID NO: 7295, SEQ ID NO: 7296, SEQ ID NO: 7297, SEQ ID NO: 7298, SEQ ID NO: 7299, SEQ ID NO: 7300 and SEQ ID NO: 7301. Preferably, the nucleotide sequence is selected from the group consisting of SEQ ID NO: 7292 and SEQ ID NO: 7293. Still more preferably, the nucleotide sequence is SEQ ID NO: 7293.

The siRNA disclosed in this application may be prepared with modified ribonucleotides as described herein. Further, the modified ribonucleotide of the siRNA used in the present method is incorporated into said siRNA by chemical synthesis or enzymatic synthesis.

The siRNA disclosed in this application may or may not have a 5' triphosphate group.

The modified siRNA is administered to a patient by a method selected from the group consisting of intravenous injection, subcutaneous injection, oral delivery, and liposome delivery. The modified siRNA accumulates in an organ, tissue or body system of the patient that are the liver, gastrointestinal tract, respiratory tract, cervix or skin.

The present invention also provides a method of inhibiting the replication of a virus, such as SARS virus, in cells positive for SARS virus comprising transfecting SARS-positive cells with a vector that directs the expression of modified siRNA which is specific for SARS. The cells are evaluated to determine if a marker in the cells has been inhibited by the modified siRNA.

The term patient, as used herein, can be an animal, preferably a mammal. More preferably the subject can be a primate, including non-human and humans. The terms subject and patient can be used interchangeably.

The treatment envisioned by the current invention can be used for subjects with a preexisting viral infection, or for subjects pre-disposed to a SARS virus infection. Additionally, the method of the current invention can be used to correct or compensate for cellular or physiological abnormalities involved in conferring susceptibility to viral infections in patients, and/or to alleviate symptoms of a viral infection in patients, or as a preventative measure in patients.

The method of treating a patient having a viral infection involves administration of compositions to the subjects. As used herein, composition can mean a pure compound, agent or substance or a mixture of two or more compounds, agents or substances. As used herein, the term agent, substance or compound is intended to mean a protein, nucleic acid, carbohydrate, lipid, polymer or a small molecule, such as a drug.

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In one embodiment of the current invention, the composition administered to the subject is a pharmaceutical composition. Further, the pharmaceutical composition can be administered orally, nasally, parenterally, intrasystemically, intraperitoneally, topically (as by drops or transdermal patch), bucally, or as an oral or nasal spray. The term "parenteral," as used herein, refers to modes of administration which include intravenous, intramuscular, intraperitoneal, intrasternal, subcutaneous and intraarticular injection and infusion. The pharmaceutical compositions as contemplated by the current invention may also include a pharmaceutically acceptable carrier.

By "pharmaceutically acceptable carrier" is intended, but not limited to, a non-toxic solid, semisolid or liquid filler, diluent, encapsulating material or formulation auxiliary of any type, such as liposomes.

A pharmaceutical composition of the present invention for parenteral injection can comprise pharmaceutically acceptable sterile aqueous or nonaqueous solutions, dispersions, suspensions or emulsions as well as sterile powders for reconstitution into sterile injectable solutions or dispersions just prior to use. Examples of suitable aqueous and nonaqueous carriers, diluents, solvents or vehicles include water, ethanol, polyols (such as glycerol, propylene glycol, polyethylene glycol, and the like), carboxymethylcellulose and suitable mixtures thereof, vegetable oils (such as olive oil), and injectable organic esters such as ethyl oleate. Proper fluidity can be maintained, for example, by the use of coating materials such as lecithin, by the maintenance of the required particle size in the case of dispersions, and by the use of surfactants.

The compositions of the present invention can also contain adjuvants such as, but not limited to, preservatives, wetting agents, emulsifying agents, and dispersing agents. Prevention of the action of microorganisms can be ensured by the inclusion of various antibacterial and antifungal agents, for example, paraben, chlorobutanol, phenol, sorbic acid, and the like. It can also be desirable to include isotonic agents such as sugars, sodium chloride, and the like. Prolonged absorption of the injectable pharmaceutical form can be brought about by the inclusion of agents which delay absorption such as aluminum monostcarate and gelatin.

In some cases, to prolong the effect of the drugs, it is desirable to slow the absorption from subcutaneous or intramuscular injection. This can be accomplished by the use of a liquid suspension of crystalline or amorphous material with poor water solubility. The rate of absorption of the drug then depends upon its rate of dissolution which, in turn, can depend upon crystal size and crystalline form. Alternatively, delayed absorption of a parenterally administered drug form is accomplished by dissolving or suspending the drug in an oil vehicle.

Injectable depot forms are made by forming microencapsule matrices of the drug in biodegradable polymers such as polylactide-polyglycolide. Depending upon the ratio of drug to polymer and the nature of the particular polymer employed, the rate of drug release can be

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controlled. Examples of other biodegradable polymers include poly(orthoesters) and poly(anhydrides). Depot injectable formulations are also prepared by entrapping the drug in liposomes or microemulsions which are compatible with body tissues.

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The injectable formulations can be sterilized, for example, by filtration through a bacterial-retaining filter, or by incorporating sterilizing agents in the form of sterile solid compositions which can be dissolved or dispersed in sterile water or other sterile injectable medium just prior to use.

Solid dosage forms for oral administration include, but are not limited to, capsules, tablets, pills, powders, and granules. In such solid dosage forms, the active compounds are mixed with at least one item pharmaceutically acceptable excipient or carrier such as sodium citrate or dicalcium phosphate and/or a) fillers or extenders such as starches, lactose, sucrose, glucose, mannitol, and silicic acid, b) binders such as, for example, carboxymethylcellulose, alginates, gelatin, polyvinylpyrrolidone, sucrose, and acacia, c) humectants such as glycerol, d) disintegrating agents such as agar-agar, calcium carbonate, potato or tapioca starch, alginic acid, certain silicates, and sodium carbonate, e) solution retarding agents such as paraffin, f) absorption accelerators such as quaternary ammonium compounds, g) wetting agents such as, for example, acetyl alcohol and glycerol monostearate, h) absorbents such as kaolin and bentonite clay, and i) lubricants such as talc, calcium stearate, magnesium stearate, solid polyethylene glycols, sodium lauryl sulfate, and mixtures thereof. In the case of capsules, tablets and pills, the dosage form can also comprise buffering agents.

Solid compositions of a similar type can also be employed as fillers in soft and hard filled gelatin capsules using such excipients as lactose or milk sugar as well as high molecular weight polyethylene glycols and the like.

The solid dosage forms of tablets, dragees, capsules, pills, and granules can be prepared with coatings and shells such as enteric coatings and other coatings well known in the pharmaceutical formulating art. They can optionally contain opacifying agents and can also be of a composition that they release the active ingredient(s) only, or preferentially, in a certain part of the intestinal tract, optionally, in a delayed manner. Examples of embedding compositions which can be used include polymeric substances and waxes.

The active compounds can also be in micro-encapsulated form, if appropriate, with one or more of the above-mentioned excipients.

Liquid dosage forms for oral administration include, but are not limited to, pharmaceutically acceptable emulsions, solutions, suspensions, syrups and elixirs. In addition to the active compounds, the liquid dosage forms can contain inert diluents commonly used in the art such as, for example, water or other solvents, solubilizing agents and emulsifiers such as ethyl alcohol, isopropyl alcohol, ethyl carbonate, ethyl acetate, benzyl alcohol, benzyl benzoate,

propylene glycol, 1,3-butylene glycol, dimethyl formamide, oils (in particular, cottonseed, groundnut, corn, germ, olive, castor, and sesame oils), glycerol, tetrahydrofurfuryl alcohol, polyethylene glycols and fatty acid esters of sorbitan, and mixtures thereof.

Besides inert diluents, the oral compositions can also include adjuvants such as wetting agents, emulsifying and suspending agents, sweetening, flavoring, and perfuming agents.

Suspensions, in addition to the active compounds, can contain suspending agents as, for example, ethoxylated isostearyl alcohols, polyoxyethylene sorbitol and sorbitan esters, microcrystalline cellulose, aluminum metahydroxide, bentonite, agar-agar, and tragacanth, and mixtures thereof.

Alternatively, the composition can be pressurized and contain a compressed gas, such as nitrogen or a liquefied gas propellant. The liquefied propellant medium and indeed the total composition is preferably such that the active ingredients do not dissolve therein to any substantial extent. The pressurized composition can also contain a surface active agent. The surface active agent can be a liquid or solid non-ionic surface active agent or can be a solid anionic surface active agent. It is preferred to use the solid anionic surface active agent in the form of a sodium salt.

The compositions of the present invention can also be administered in the form of liposomes. As is known in the art, liposomes are generally derived from phospholipids or other lipid substances. Liposomes are formed by mono- or multi-lamellar hydrated liquid crystals that are dispersed in an aqueous medium. Any non-toxic, physiologically acceptable and metabolizable lipid capable of forming liposomes can be used. The present compositions in liposome form can contain, in addition to the compounds of the invention, stabilizers, preservatives, excipients, and the like. The preferred lipids are the phospholipids and the phosphatidyl cholines (lecithins), both natural and synthetic. Methods to form liposomes are known in the art (see, for example, Prescott, Ed., Meth. Cell Biol. 14:33 et seq (1976)).

One of ordinary skill will appreciate that effective amounts of the agents of the invention can be determined empirically and can be employed in pure form or, where such forms exist, in pharmaceutically acceptable salt, ester or prodrug form. The agents can be administered to a subject, in need of treatment of viral infection, as pharmaceutical compositions in combination with one or more pharmaceutically acceptable excipients. It will be understood that, when administered to a human patient, the total daily usage of the agents or composition of the present invention will be decided by the attending physician within the scope of sound medical judgement. The specific therapeutically effective dose level for any particular patient will depend upon a variety of factors: the type and degree of the cellular or physiological response to be achieved; activity of the specific agent or composition employed; the specific agents or composition employed; the age, body weight, general health, sex and diet of the patient; the time

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of administration, route of administration, and rate of excretion of the agent; the duration of the treatment; drugs used in combination or coincidental with the specific agent; and like factors well known in the medical arts. For example, it is well within the skill of the art to start doses of the agents at levels lower than those required to achieve the desired therapeutic effect and to gradually increase the dosages until the desired effect is achieved.

Dosing can also be arranged in a patient specific manner to provide a predetermined concentration of the agents in the blood, as determined by techniques accepted and routine in the art. Thus patient dosaging can be adjusted to achieve regular on-going blood levels, as measured by HPLC, on the order of from 50 to 1000 ng/ml.

It will be readily apparent to one of ordinary skill in the relevant arts that other suitable modifications and adaptations to the methods and applications described herein can be made without departing from the scope of the invention or any embodiment thereof.

The modified siRNA is prepared by custom chemical synthesis by Dharmacon, at Lafayette CO. Each C and U within the siRNA duplex (GL2), has been substituted with 2'-F-U and 2'-F-C except for the 3'-end overhang, which was dTdT.

To test the stability of 2' chemically modified siRNA compared to unmodified siRNA (siRNA), the following experiment is performed. 4ngs of siRNA are added to a 20 μ L volume of 80% human serum from a healthy donor. This mixture is incubated at 37°C for various times ranging from 1 minute up to 10 days. The same process is performed for 2' fluorine modified siRNA (2'-F siRNA). When the incubation process is finished, the mixtures are placed on ice and then immediately separated by PAGE along with a 32 P-siRNA control. The 2' modified siRNA is stable as compared to unmodified siRNA.

V. IDENTIFICATION OF THERAPEUTICALLY ACTIVE AGENTS FOR TREATMENT OF SARS VIRUS INFECTION

The invention provides methods for treating SARS by administering therapeutically active agents, such as small molecule compounds, to a mammal, as well as methods of identifying therapeutically active agents, such as potent small molecules, for the treatment of SARS virus infection.

In one aspect of the invention a method of identifying a therapeutically active agent is provided comprising: (a) contacting the therapeutically active agent with a cell infected with the SARS virus; (b) measuring attenuation of a SARS related enzyme.

In a more particular embodiment, the therapeutically active agent is a small molecule. In another more particular embodiment, the therapeutically active agent is a nucleoside analog (e.g. Ribavirin). In another more particular embodiment the small molecule is a SMIP or peptidic immunomodulating compound. In another more particular embodiment the therapeutically

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active agent is a peptoid, oligopeptide, or polypeptide. In another embodiment the SARS related enzyme is SARS protease. In another embodiment the SARS related enzyme is SARS polymerase. In still another embodiment the SARS related enzyme is a kinase. In still another embodiment, the SARS related enzyme is a protease. The furin inhibitor peptidyl chloromethylketone prevents blocks cell-cell fusion after MHV infection (de Haan et al. (2004) J Virol), which offers guidance for SARS therapy.

The invention includes a cell-based assay that can be used to screen for and identify a therapeutically active agent for the treatment of SARS virus infection. Therapeutically active agents of the invention include agents that inhibit, prevent or reduce the replication of a SARS virus. Such agents can be identified by infecting a cultured cell (such as, for example, VERO cells) with a SARS virus and evaluating the impact of potential antiviral compounds on SARS virus replication. Assays to measure the effect of a potential antiviral compound on virus replication are known in the art and may be based on a variety of parameters.

The cell-based assay may be used in a high-throughput screen to identify therapeutically active compounds from chemical libraries comprising potential antiviral compounds.

Therapeutically active compounds suitable for use in the invention may inhibit any SARS viral target that is essential for viral replication in whole cells. Efficacy (the ability of a compound to inhibit or inactivate the target, be it viral or cellular, that results in the reduction of virus in the culture) of the therapeutic agent is measured by assessing the viability and/or the proliferation of surviving cells in a SARS virus infected cell culture.

A number of methods can be used to measure cell viability are known in the art, such as assays measuring cellular enzymes, proteins, nucleotide triphoshates (e.g. ATP), nucleic acids (e.g. host cell mRNA (e.g. GAPDH) or rRNA sequences) or cellular metabolites such as MTT or MTS. In addition, fluorescent (including, for example HSV paper) or non-fluorescent dyes (e.g. propidium diiodide) or labeling of DNA can be used to measure indications of cell viability and/or proliferation.

Alternatively, efficacy of a compound or sample can be determined by directly measuring the amount of virus or viral products in the culture. Methods for measuring the amount of virus, viral genome or viral products include: PCR, RT-PCR, TMA, reporter proteins with fluourescent or luminescent qualities or enzymatic functions (e.g., luciferase, alkaline phosphatase, GFP) or proteins that can be detected by antibodies (e.g. EGF) that might be incorporated into the viral genome prior to infection of the cell culture. Further, viral products such as viral proteins can be measured by ELISA or enzymatic activities. Methods for identifying viral polynucleotides, viral proteins and antibodies specific to viral proteins are discussed above.

Potential antiviral compounds are applied to the cell-based assay at a concentration of approximately 10 μ M and compound classes having therapeutic effect are identified by

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measuring the parameter of choice (such as cell viability/proliferation or the virus or viral genome or a viral product be it viral in origin or non-virus in origin). Once compounds are identified as having activity, they are resynthesized, and analoged. Starting with the identified compound, many analogs and new compounds are synthesized during consecutive optimization cycles of synthesis, biological profiling and modeling techniques to optimize the to the lead structure until *in vivo* activity is elucidated and optimized.

Cells suitable for use in the assay include the cells described above as suitable for vaccine production. Preferably, the cells are African green monkey kidney cells (Vero) cells. Human embyronic lung fibroblasts or normal human diploid fibroblasts may also be used in the invention.

. In one embodiment, the invention includes a fluorescence based cytopathogenicity assay to measure the effect of a potential antiviral compound on a cell-based assay. One example of a fluorescence based cytopathogenicity assay is illustrated below.

 1×10^4 Vero cells per well of a microtiter plate (MTP) are infected with a defined amount of SARS virus selected within the following ranges for optimal MOI: 5-10, 10-25, 25-50, 50-100, 100-500, or 500-1000 PFU in a total volume of 200 μ l media (M199 medium supplemented with 5% FCS, 2 mM glutamine, 100 IU/ml penicillin and 100 μ g/ml streptomycin) in the presence or absence of the potential antiviral compound and incubated for at least 1, 2, 3, 4, 5, 6, or 7 days at 37°C, 5% CO₂. The wells of the MTP are washed with PBS (200 μ l) and then filled with 200 μ l PBS containing 10 μ g/ml fluorescein diacetate. After a 45-min incubation at room temperature, fluorescence is measured at 485 nm excitation and 538 nm emission wavelengths. IC₅₀ values are determined by a nonlinear plot of antiviral activity as a function of drug concentration.

Other cell based assays are known in the art and include, among others, methods of GFP detection and Luc detection. In addition, a Promega kit is commercially available that provides additional methods of measuring cell viability, etc.

In one embodiment, the invention includes a method of measuring the efficacy of a potential antiviral compound using RT-PCR to detect the levels of SARS viral RNA in the cell based assay. Methods of using RT-PCR are known in the art. One example of such an assay is described below.

 5×10^6 Vero cells are seeded in tissue culture. Flasks containing the cells are incubated over night at 37°C, 5% CO₂. The cells are infected (m.o.i. = 1) with SARS virus in the presence and absence of potential antiviral compounds. Optionally, the cells may be pretreated with the potential compound prior to infection. In either case, a suitable control cell assay is also prepared.

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The RNA of infected cells is purified at 2 h (UL54), 12 h (UL8) and 16 h (UL13) after infection, (Qiagen) RNA purification (RNeasy kit; 40 μ l elution) and quantified (absorption at 260 nm). The RNA (2 μ g) is reverse transcribed with a specific primer (2 pmol, using one of the primer pairs described herein) into cDNA according to the Superscript II protocol (Invitrogen). Aliquots (2 μ l) of the reverse transcription reaction are amplified by PCR. Fragments of the appropriate target SARS gene, i.e., a gene encoding a SARS enzyme, are amplified in 30 cycles (UL54 and UL8: 3 min, 94°C hot start; 1 min, 94°C denaturation; 1 min, 55°C annealing; 1 min, 72°C polymerization. UL13: 3 min, 94°C hot start; 1 min, 94°C denaturation; 1 min, 60°C annealing; 1 min, 72°C polymerization) by PCR (Taq-Polymerase, Stratagene), in a 100- μ l reaction volume with the appropriate oligonucleotides, as described herein at 0.1 nmol each. 8- μ l aliquots of cycle 20–30 (lanes 2–12) of the PCR were resolved on a 2% agarose gel (Invitrogen) according to the manufacturer's instructions.

Cell-based assays of the invention may optionally use a variant or derivative of a wild-type SARS virus that has reduced or attenuated virulence in humans and/or animal models (e.g., mouse, non-human primate, etc.) Use of such attenuated SARS viruses in screening methods may reduce safety concerns and precautions that would otherwise be associated with the pathogenic nature of the SARS virus and may eliminate or reduce the need for the implementation of cumbersome high containment levels during performance of the assays and screening of compounds.

The invention includes an enzyme-based assay that can be used to screen for and identify a therapeutically active agent for the treatment of SARS virus infection.

An embodiment of the invention is an assay comprising contacting a known quantity of SARS protease in solution to a peptide containing a detectable marker and cleavage site for SARS protease, wherein SARS protease activity is monitored by measuring the intensity of the marker on the cleaved product.

In a more particular embodiment, a method of assaying for SARS protease is provided comprising contacting a sample solution containing SARS protease with a peptide containing a fluorescent donor, fluorescent quencher, and cleavage site for SARS protease, said peptide being detectable with a fluorometer when cleaved, wherein SARS protease activity is determined in the sample by the amount of fluorescence detected by the fluorometer.

Assays based on the direct measurement of SARS protease inhibition may be utilized for screening for SARS therapeutics. Protease for such assays such as 3C-like protease and papain-like protease may be isolated and purified for such assays as described in Seybert, et al., J. Gen. Virol., 78:71-75, 1997, Ziebuhr, et al., Adv. Exp. Med. Biol., 440:115-120, 1998, Sims, et al., Adv. Exp. Med. Biol. 440:129-134, 1998, Ziebuhr, et al., J. Virol., 73:177-185, 1999, Teng, et al., J. Virol., 73:2658-2666, 1999, Herold, et al., J. Biol. Chem. 274:14918-14925, 1999, and

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Ziebuhr, et al., J. Biol. Chem. 276:33220-33232, 2001. Furthermore, Example 30 describes a novel method of purifying SARS protease using column chromatography. Example 31 describes a continuous fluorescence resonance energy transfer (FRET) assay for measuring SARS protease activity. Protease enzyme based assays such as the FRET assay demonstrated in Example 31 are readily adapted for high-throughput screening and are used for screening candidate antiviral compounds. Performance of the protease enzymatic assay in the presence of a SARS protease inhibitor compound will show a decreased amount of fluorescence at a given time when compared to negative control assay containing no test compound on a non-inhibiting control compound. Such a method would involve the steps of: (a) providing an assay solution comprising SARS protease; (b) adding a test compound to the assay solution; (c) adding a substrate for SARS protease to the assay solution; and (d) measuring the proteolytic activity in the assay solution. In a preferred embodiment, the proteolytic activity is measured by the fluorescence of fluorophore product produced by the enzymatic activity of SARS protease.

Attenuated SARS virus variants generally contain one or more genome modifications or mutations (e.g., substitutions, deletions, insertions) in protein encoding or non-coding regions. Specific examples of attenuating mutations include, for example, genetic modifications in the 5'-end noncoding region, leader sequence, intergenic regions, 3'-end noncoding region, ORF 1a, ORF 1b, S gene, E gene, M gene, N gene, or any of the nonstructural protein genes outside of the ORF 1a/1b region. Preferred attenuating mutations are in a SARS virus structural protein (e.g., Spike (S)), a protease or polymerase domain, or a non-coding sequence (e.g., 5'-end noncoding region, intergenic sequence). In addition, a cleavage site may be introduced or eliminated within the spike protein (see for example, Gombold et al., J. Virol. 67:4504-4512, 1993; Bos et al., Virology 214:453-463, 1995), such modification that may also be useful for optimization of expression of recombinant spike protein antigen (e.g., for vaccine purposes).

A variety of methods are used according to the present invention in order to obtain attenuated variants of SARS virus. Such methods include serial passage of the SARS virus in cultured cells (e.g., mammalian cell culture, such as fetal rhesus kidney cells or VERO cells), until the SARS virus demonstrates attenuated function. The serial propagation of virus may be performed at any temperature at which tissue culture passage attenuation occurs, and may be performed in conjunction with one or more steps of mutagenesis (e.g., chemical mutagenesis). The attenuated phenotype of SARS virus variants, obtained after one or more cell culture passages, is readily measured by one skilled in the art. As used herein, attenuated refers to the decreased virulence of the SARS virus in a human subject. Evidence of attenuated function may be indicated by decreased levels of viral replication or by decreased virulence in an animal model.

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Other methods of producing an attenuated SARS virus include cell culture passage of the virus at sub-optimal temperatures (cold passage), as well as introduction of attenuating mutations into the SARS viral genome by random mutagenesis (e.g., chemical mutagenesis, such as using 5-fluorouracil) or using directed mutagenesis. Preparation and generation of attenuated RSV vaccines (the methods of which will generally applicable to SARS virus) are disclosed in, for example, EP 0640128, US Patent No. 6,284,254, US Patent No. 5,922,326, US Patent No. 5,882,651.

The number of passages required to obtain safe, immunizing attenuated virus is dependent at least in part on the conditions employed. Periodic testing of the SARS virus culture for virulence and immunizing ability in animals (e.g., mouse, primate) can readily determine the parameters for a particular combination of tissue culture and temperature.

In another embodiment, the cell-based assay for screening of antiviral compounds is based on the readout of expression of a gene product (e.g., reporter gene product) that is not from SARS virus. Gene products particularly suitable to the present invention include, but are not limited to those of the above-described assays.

In order to achieve such a read-out, the gene-of-interest (GOI) encoding said gene reporter gene product must be incorporated into a replicating SARS virus genome or construct derived from a SARS virus genome (e.g., SARS virus replicon, SARS virus defective-interfering (DI) RNA). Figure 13 is a schematic depicting locations for incorporation of the reporter gene into a SARS virus genome. Preferably, insertion of a heterologous reporter gene-of-interest is at a site between existing SARS virus genes, such as for example, as shown in Figure 13. For example, the GOI may be inserted closely following the termination codon of a SARS virus gene (e.g., ORF 1b, S, E, M, N). Insertion should be positioned in order to minimize disruption of mRNA transcription for the SARS virus gene(s). The GOI may also be inserted as an in-frame "fusion" with an existing SARS virus gene, such that sufficient function of the GOI is maintained for detection. To optimize expression, an additional SARS virus intergenic sequence (e.g., SEQ ID NO: 7388, with or without additional flanking SARS virus sequences) may also be engineered into a position preceding the inserted GOI.

Incorporation of a GOI into SARS virus may be accomplished by one of skill in the art using a variety of techniques. For example, one preferred method is targeted RNA recombination, that takes advantage of the ability of coronavirus RNAs to undergo recombination within the cell (see for example Fischer et al., J. Virol. 71:5148-5160, 1997; Koljesar et al., J. Vet. Sci. 2:149-157, 2001). A construct of desired configuration (e.g., cDNA of defective interfering RNA of SARS virus) containing the GOI flanked by SARS virus sequence (e.g., intergenic sequence) is generated such that RNA may be transcribed directly within a eukaryotic cell or in vitro and transfected into susceptible cells also infected with SARS

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virus. Recombinant virus containing the GOI is identified based on expression of the GOI encoded marker.

Alternatively, incorporation of a GOI into SARS virus may be accomplished by one of skill in the art by first assembling a full-length cDNA clone of the SARS virus, that can be used to produce infectious RNA transcripts in vivo (e.g., from an RNA polymerase II promoter) or in vitro (e.g., from a bacteriophage promoter). Although relatively long in genome length, such assembly of a full-length cDNA clone is now readily obtainable by one of skill in the art using standard molecular biology and reverse genetics techniques and the genome sequence of SARS virus (see for example, Thiel et al., J. Gen. Virol., 82:1273-1281, 2001; Almazan et al., Proc. Natl. Acad. Sci. USA 97:5516-5521, 2000; Thiel et al. (2003) J Gen Virol 82:1273-1281; Yount et al (2003) PNAS USA 100:12995-13000). Insertion of a heterologous GOI into a full-length SARS virus genome cDNA may be performed using a variety of techniques, such as for example, ligation into natural or synthetic restriction sites, PCR (e.g., overlapping PCR), and recombination.

It may also be desirable to utilize similar SARS virus recombinants containing a gene-ofinterest for antiviral screening, however, with further modification to minimize or eliminate virus-induced cytopathology (e.g., CPE). Non-cytopathic derivatives from SARS virus may be obtained by one of skill in the art using a variety of methods. For example, a selectable marker (e.g., drug resistance marker) may be incorporated as GOI into a SARS virus genome to produce infectious virus as described above (see for example, Perri et al., J. Virol., 74:9802-9807, 2000). Infectious GOI-containing SARS virus or infectious genome RNA/cDNA is then used to infect/transfect cells (e.g., VERO), with or without prior mutagenesis, after which time the infected cells are subjected to the appropriate selection. Only those cells containing SARS virus harboring both the selectable marker and one or more mutations rendering the virus noncytopathic will survive the selection process and grow out. Active SARS virus replication in these cells is readily detected using a variety of detection techniques (e.g., PCR, Northern blot) and such cells may serve as the substrate for cell-based screening assays. Mutations that result in the desired noncytopathic SARS virus phenotype may include nucleotide substitutions, deletions or additions, and may occur in a variety of genome coding or non-coding regions (e.g., 5' or 3'end noncoding regions, intergenic regions, ORF1a, ORF1b, a protease domain, a polymerase domain). The identification of such mutations is readily accomplished by exchange of sequences with wild-type (e.g., parental) SARS virus and demonstrating transfer of the phenotype, and sequencing of the appropriate genome region. Similar mutations that reduce or eliminate cytopathogenicity also may be utilized in the context of a SARS virus derived replicon vector, either by similar selection directly using a SARS virus replicon or by specific engineering of the replicon based on mutation(s) identified in the context of infectious SARS virus as described

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above. In addition, such mutations may serve as the basis for attenuated SARS virus derivatives, as described elsewhere in this document.

Alternatively, rather than using infectious SARS virus or its derivatives for cell-based screening assays, propagation defective "replicons" may be engineered and utilized. Such replicons maintain all protein encoding sequences and cis replication sequences required for RNA replication and expression within a cell, but are deleted of one or more sequences or genes required for packaging of progeny SARS virus (see for example Curtis et al., J. Virol., 76:1422-1434, 2002). Figure 14 is a schematic depicting representative examples of SARS virus replicons according to the present invention. For example a SARS virus cDNA construct is generated, that is lacking one or more (or all) structural protein encoding genes, whereby the missing SARS virus gene(s) is/are replaced by the GOI, maintaining all necessary transcription signals for expression of the GOI. Operably linked with the SARS virus replicon cDNA construct is a promoter for RNA polymerase that can be used to transcribe the replicon RNA in vivo (e.g., RNA polymerase II promoter) or in vitro (e.g., bacteriophage promoter). The SARS replicon may be introduced into a susceptible cell by transfection as RNA or DNA, depending on the promoter of choice, and the transfected cells may be utilized for the evaluation of antiviral compounds. By incorporating one or more mutations rendering the replicon noncytopathic for the cells (see above), one can avoid the need for nucleic acid transfection each time an assay is to be performed.

Alternatively, SARS virus replicons may be packaged into virus like particles that allow infection of cells, rather than requiring transfection of nucleic acid molecules. A requirement for replicon packaging is that essential SARS virus gene functions deleted from the replicon (e.g., one or more structural proteins) are provided in trans within the cell containing the replicon. A variety of methods for packaging of replicon RNA can be utilized to one of skill in the art (see for example, Curtis et al., ibid: Ortego, et al., J. Virol., 76:11518-11529, 2002). For example, stably transformed cell lines constitutively or inducibly expressing the required SARS virus gene functions may be utilized. Alternatively, the required SARS virus gene functions may be expressed by viral vectors that are introduced into the replicon-containing cell. Alternatively a defective interfering (DI) SARS virus derived RNA containing the required gene functions may be introduced into the replicon-containing cell. Such DI constructs used to complement missing replicon functions may be more commonly referred to as defective helper RNA or defective helpers.

Another configuration useful for cell-based antiviral screening assays according to the present invention utilizes SARS virus derived DI RNAs encoding a GOI (see for example Stirrups, et al., J. Gen. Virol., 81:1687-1698, 2000; Liao, et al., Virology 208:319-327, 1995).

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Introduction of a SARS DI, either as cDNA linked to an RNA polymerase II promoter or as in vitro transcribed RNA, into susceptible cells also infected with SARS virus, allows for a readout of the GOI reporter product in assays.

A replicon-based system for rapid identification of coronavirus replicase inhibitors is described by Hertzig et al. (2004) J Gen Virol DOI 10.1099/vir/0/80044-0. Briefly, the system uses a non-cytopathic selectable replicon RNA that can be stably maintained in eukaryotic cells. The replicon RNA mediates reporter gene expression as a marker for coronavirus replication, and expression of the reporter can be used to test the inhibitory effect of test compounds in vitro, thereby allowing high throughput screening for replicase inhibitors without the need to grow infectious virus. Preferred replicon RNAs include a neomycin resistance gene in the replicase gene with a downstream reporter gene (e.g. GFP) that is expressed via replicase-mediated synthesis of a sub-genomic mRNA.

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VI. COMPOSITIONS AND METHODS FOR TREATMENT OF SARS VIRUS INFECTION The present invention relates to compositions and methods for the treatment and/or prevention of SARS. The invention further includes a method for the treatment and/or prevention of SARS through the administration of a therapeutically effective amount of at least one antiviral compound from among those described in the US Patents and published international patent applications listed in Table 1 and Table 2. In one embodiment of the method, the antiviral compound is a small molecule. In another embodiment, the antiviral compound is a protease inhibitor. In a further embodiment, the antiviral protease inhibitor is a 3C-like protease inhibitor. and/or a papain-like protease inhibitor. Combined treatment with the lopinavir/ritonavir (Kaletra) protease inhibitor and ribavirin has shown a favorable clinical response (Chu et al. (2004) Thorax 59:252-256). In another embodiment, the antiviral compound is an inhibitor of an RNA dependent RNA polymerase. In another embodiment, a first antiviral compound that is a protease inhibitor is administered with a second antiviral compound that is an RNA-dependent RNA polymerase inhibitor. The invention further provides for the administration of a steroidal anti-inflammatory drug in combination with at least one antiviral compound, for example, from the antiviral compounds described in the documents listed in Table 1 and Table 2. A combination treatment of steroids and ribavirin has been described by Fujii et al. (2004) J Infect Chemother 10:1-7. A combination treatment of corticosteroids and interferon alfacon-1 has also been reported (Loutfy et al. (2003) JAMA 290:3222-3228).

The invention further provides for a method for the treatment and/or prevention of SARS through the administration of a therapeutically effective amount of at least one antiviral

compound from among those described in the US Patents and published international patent applications listed in Table 1 and Table 2 by inhalation. In another aspect, the antiviral compound may be administered in combination with a SMIP, SMIS, or other immunomodulatory compound such as those in Table 34 and in Table 35. In one embodiment of the method, the antiviral compound is a small molecule. In another embodiment, the antiviral compound is a protease inhibitor. In a further embodiment, the antiviral protease inhibitor is a 3C-like protease inhibitor and/or a papain-like protease inhibitor. In another embodiment, the antiviral compound is an inhibitor of an RNA dependent RNA polymerase. In another embodiment, a first antiviral compound that is a protease inhibitor is administered with a second antiviral compound that is an RNA-dependent RNA polymerase inhibitor. The invention further provides for the administration of a steroidal anti-inflammatory drug in combination with at least one antiviral compound, for example, from the antiviral compounds described in the documents listed in Table 1 and Table 2. The steroidal anti-inflammatory drug may be administered by inhalation for a local effect or administered for systemic absorption such as via an oral or intravenous route.

The invention further provides for methods for treating SARS infection comprising administering a small molecule immunopotentiator (SMIP) compound either alone or in combination with an antiviral compound or in combination with a SARS vaccine. In a further embodiment, the SMIP is a compound disclosed herein or set forth in Table 34.

The invention further provides for methods for treating SARS infection comprising administering an immunosuppressant compound, optionally a small molecule suppressant (SMIS) compound either alone or in combination with an antiviral compound. In a further embodiment, the immunosuppressant compound is disclosed herein or set forth in Table 35.

The invention further provides peptidic immunomodulating compositions, that include

oligo and polypeptides, capable of effecting inflammatory response in a patient. In one embodiment, the peptidic immunomodulating composition is able to stimulate human cells to produce cytokines. In another embodiment the peptidic immunomodulating composition is capable of decreasing cytokine levels in the human. Preferred Examples of peptidic immunomodulating compositions include those listed in Table 35, as well as TGF β 2, TGF β 1, TGF β 3, thymopentin (TP5), β -mercaptopropionyl-arginyl—lysyl-aspartyl-valyl-tyrosyl-cysteine amide, colostrinine, lactoferrin (LF), cyclolinopeptide A (CLA), and tuftsin (TKPR). The peptidic immunomodulating compositions of the invention may be used alone or in combination

with other agents, preferably antiviral compounds, for the treatment of SARS.

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The invention further provides for a kit for use by a consumer for the treatment and/or prevention of SARS. Such a kit comprises: a) a pharmaceutical composition comprising a therapeutically effective amount of at least one antiviral, SMIP, SMIS, or other immunomodulating compound from among those described in the US Patents and published international patent applications listed in Table 1, Table 2, Table 34 and Table 35 and a pharmaceutically acceptable carrier, vehicle or diluent; b) a container for holding the pharmaceutical composition; and, optionally, c) instructions describing a method of using the pharmaceutical compositions for the treatment and or the prevention of SARS. The kit may optionally contain a plurality of compounds for the treatment of SARS wherein the antiviral compounds are selected from 3C-like protease inhibitors and papain-like protease inhibitors. In a further embodiment, the kit contains an antiviral compound that is an RNA-dependent RNA polymerase inhibitor. When the kit comprises more than one antiviral, SMIP, SMIS, or other immunomodulating compound, the compounds contained in the kit may be optionally combined in the same pharmaceutical composition.

An additional aspect of the invention provides for the use of at least one of the antiviral, SMIP, SMIS, or other immunomodulating compounds described in the US Patents and published international patent applications listed in Table 1, Table 2, Table 34 and Table 35 for the manufacture of a medicament for the treatment or prevention of SARS.

An additional aspect of the invention provides for the use of at least one SMIP compound, or at least one immunosuppressant compound, or at least one SMIS compound for the manufacture of a medicament for the treatment or prevention of SARS. Preferred SMIP, immunosuppressant, and SMIS compounds are described herein.

Unless otherwise specified, the following terms, when used within Section VI: "Compositions and Methods for Treatment of SARS Virus Infection" of the present application have the meanings as defined below:

As used herein, "limit", "treat" and "treatment" are interchangeable terms as are "limiting" and "treating" and, as used herein, include preventative (e.g., prophylactic) and palliative treatment or the act of providing preventative or palliative treatment. The terms include a postponement of development of SARS symptoms and/or a reduction in the severity of such symptoms that will or are expected to develop following infection with a SARS virus. The terms further include ameliorating existing SARS symptoms, preventing additional symptoms, ameliorating or preventing the underlying metabolic causes of symptoms.

Representative uses of the compositions and methods of the present invention include: the elimination or reduction of the viral load of the SARS virus in a vertebrate, including humans, the elimination or reduction of symptoms associated with SARS, and a reduction in morbidity

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associated with SARS. In a SARS patient population, the use of the compositions and methods of the invention will result in the reduction in the high mortality rates associated with SARS.

Infection with the SARS virus and the symptoms associated with SARS can be treated in a subject by administering the compositions of the invention. The compositions of the invention may be administered systemically. For systemic use, the compounds herein are formulated for parenteral (e.g., intravenous, subcutaneous, intramuscular, intraperitoneal, intranasal or transdermal) or enteral (e.g., oral or rectal) delivery according to conventional methods.

Intravenous administration can be by a series of injections or by continuous infusion over an extended period. Administration by injection or other routes of discretely spaced administration can be performed at intervals ranging from weekly to once to three times daily or more.

Alternatively, the compositions disclosed herein may be administration, followed by administration of disclosed compositions, followed by no administration, followed by administration of disclosed compositions, and the like). Treatment will continue until the desired outcome is achieved.

A "subject" is a vertebrate animal including a human that is in need of treatment with the compositions, methods and kits of the present invention. The term "subject" or "subjects" is intended to refer to both the male and female gender unless one gender is specifically indicated.

"Coadministration" of a combination of a plurality of antiviral compounds means that these components can be administered together as a composition or as part of the same, unitary dosage form. "Co-administration" also includes administering a plurality of antiviral compounds separately but as part of the same therapeutic treatment program or regimen. "Co-administration" also includes administering a plurality of other agents, such as, for example an oligopeptide, a polypeptide, a peptidic immunomodulator, nucleic acid, antibodies, or a vaccine wherein the compounds or agents are administered separately but as part of the same therapeutic treatment program or regimen. The components need not necessarily be administered at essentially the same time, although they can if so desired. "Co-administration" also includes separate administration at different times and in any order. For example, where appropriate a patient may take one or more component(s) of the treatment in the morning and the one or more of the other component(s) at night.

By "antiviral compound" as used herein is meant an antiviral compound as described in the US Patents and published international patent applications listed in Table 1 and Table 2. The US Patents and published international patent applications listed in Table 1, Table 2 and Table 35 are incorporated herein in their entirety. In one embodiment, the antiviral compound is an RNA-dependent RNA polymerase. In another preferred embodiment the antiviral compound is a 3C-like protease inhibitor or a papain-like protease inhibitor. The antiviral compounds may be

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administered in the form of the acid, or of a soluble alkali metal salt or alkaline earth metal salt where appropriate.

The precise dosage of the antiviral compound will vary with the dosing schedule, the oral potency of the particular antiviral compound chosen, the age, size, sex and condition of the subject, the severity of the disorder to be treated, and other relevant medical and physical factors. Thus, a precise pharmaceutically effective amount cannot be specified in advance and can be readily determined by the caregiver or clinician.

Generally, an appropriate amount of antiviral compound is chosen to obtain a reduction in the load of the SARS virus in the subject and/or to obtain a reduction in the symptoms associated with SARS. For humans, an effective oral dose of antiviral compound is typically from about 1.5 to about 6000 μ g/kg body weight per day and preferably about 10 to about 2000 μ g/kg of body weight per day.

One of ordinary skill in the art will recognize that certain antiviral, SMIP, SMIS, and immunomodulating compounds of the invention including 3C-like protease inhibitors, papain-like protease inhibitators, and RNA-dependent RNA polymerase inhibitors will contain one or more atoms that may be in a particular stereochemical, tautomeric, or geometric configuration, giving rise to stereoisomers, tautomers and configurational isomers. All such isomers and mixtures thereof are included in this invention, when active. Crystalline and amorphous forms of the antiviral compounds of this invention are also included as are hydrates, solvates, polymorphs, and isomorphs of the antiviral compounds of the invention.

SMIP compounds of the invention include compounds are described in issued U.S. Patent Nos. 4,547,511 and 4,738,971 with the general structure (a):

for the treatment of disorders responsive to agents that enhance cell-mediated immunity.

Immunostimulatory oligonucleotides and polynucleotides are described in PCT WO 98/55495 and PCT WO 98/16247. U.S. Patent Application No. 2002/0164341 describes adjuvants including an unmethylated CpG dinucleotide (CpG ODN) and a non-nucleic acid adjuvant. U.S. Patent Application No. 2002/0197269 describes compositions comprising an antigen, an antigenic CpG-ODN and a polycationic polymer.

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Additionally, issued U.S. Patent Nos. 4,689,338, 5,389,640, 5,268,376, 4,929,624, 5,266,575, 5,352,784, 5,494,916, 5,482,936, 5,346,905, 5,395,937, 5,238,944, 5,525,612, WO99/29693 and U.S. Ser. No. 09/361,544 disclose compounds of the general structure (b):

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for the use as "immune response modifiers."

Further compounds with SMIP and antiviral activity are described below and in US

Patent Application entitled Thiosemicarbazones as Anti-Virals and Immunopotentiators filed on

December 29, 2003 with an attorney docket number of PP19814.004US generally disclosing
compounds of the following structures:

A compound of formula c:

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wherein: E is absent or selected from the group consisting of alkyl, substituted alkyl, cycloalkyl, substituted cycloalkyl, aryl, substituted aryl, heterocyclyl, substituted heterocyclyl, heteroaryl, and substituted heteroaryl;

L is absent or is selected from the group consisting of oxo, amino, alkylene, substituted alkylene, alkoxy, alkylamino, aminoalkyl, heterocyclyl, carbocyclyl, and carbonyl:

W is absent or selected from the group consisting of cycloalkyl, substituted cycloalkyl, aryl, substituted aryl, heterocyclyl, substituted heterocyclyl, heteroaryl, and substituted heteroaryl;

X is absent or is selected from the group consisting of oxo, amino, alkylene, substituted alkylene, alkoxy, alkylamino, aminoalkyl, heterocyclyl, carbocyclyl, and carbonyl;

Y is selected from the group consisting of cycloalkyl, substituted cycloalkyl, aryl, substituted aryl, heterocyclyl, substituted heterocyclyl, heteroaryl, and substituted heteroaryl;

Y' is absent or is selected from the group consisting of F, Cl, Br, I, nitro, alkyl, substituted alkyl, and optionally substituted heterocyclyl, amino, alkylamino, dialkylamino; Y" is absent or is selected from the group consisting of F, Cl, Br, I, nitro, alkyl, substituted alkyl, and optionally substituted heterocyclyl, amino, alkylamino, dialkylamino;

5 R' is H, alkyl, or substituted alkyl;

R" is H. or

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R' and R'' are taken together to form a hetercyclic ring;
Z and Z' are independently selected from the group consisting of hydrogen, alkyl, substituted alkyl, aryl, substituted aryl, arylalkyl, substituted arylalkyl, heteroaryl, substituted heteroaryl, heteroarylalkyl, substituted heteroarylalkyl, alkoxy, substituted alkoxy, aminocarbonyl, alkoxycarbonyl, carboxyl sulfonyl, methanesulfonyl, and substituted or unsubstituted alkylcarbonyl, arylcarbonyl, aralkylcarbonyl, heteroarylcarbonyl, heteroaralkylcarbonyl, alkylcarbonyloxy, arylcarbonyloxy, aralkylcarbonyloxy, heteroarylcarbonyloxy, heteroaralkylcarbonyloxy, alkylcarbonyloxy, arylaminocarbonyloxy, formyl, loweralkylcarbonyl, loweralkoxycarbonyl, aminocarbonyl, aminoaryl, alkylsulfonyl,

loweralkylcarbonyl, loweralkoxycarbonyl, aminocarbonyl, aminoaryl, alkylsulfonyl, sulfonamido, aminoalkoxy, alkylamino, heteroarylamino, alkylcarbonylamino, alkylaminocarbonylamino, arylaminocarbonylamino, aralkylcarbonylamino, heteroarylcarbonylamino, arylcarbonylamino, cycloamidino, cycloalkyl, cycloimido, arylsulfonyl and arylsulfonamido; or

Z and Z' are taken together to form a heterocyclic group, that may be optionally substituted and the tautomers and the pharmaceutically acceptable salts, esters, or prodrugs thereof.

Further SMIP compounds are described below and in US Patent Application 10/762873, Use of Tryptanthrin Compounds for Immune Potentiation, filed on January 21, 2004 and disclosing the general embodiment of compounds represented by Formula (d):

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wherein

A, B, C, D, E, F, G, and H are independently selected from carbon and nitrogen, or A and B and/or C and D can be taken together to be nitrogen or sulfur;

 R_1 , R_2 , R_3 , R_4 , R_8 , and R_{10} are independently selected from the group consisting of hydrogen, halogen, loweralkyl, alkyl, substituted alkyl, cycloalkyl, heterocyclyl, alkylheterocyclyl, substituted heterocyclyl, substituted alkenyl, amino, (substituted alkyl)(alkyl)amino, imino, haloloweralkyl, hydroxy, alkoxy, substituted alkoxy, hydroxyalkylthio, nitro, alkylsulfonyl, N-alkylsulfonamide, arylalkyl, arylalkylaryl, arylaryl, aryloxy, arylamino, acylamino, acylamino, alkylaminoacylamino, alkylaminosulfonylamino, alkylamino, alkenylamino, dialkylamino, alkoxyalkylamino, alkoxyalkylheterocyclyl, mercaptoalkoxyalkyl, cyano, formyl, -COOR_{11} wherein R_{11} is hydrogen, loweralkyl, aryl, heterocyclyl, monosaccharide or disaccharide, and -CONR_{12}R_{13} wherein R_{12} and R_{13} are independently selected from hydrogen, loweralkyl, aryl, heterocyclyl, saccharide, peptide and amino acid residues; or R_2 and R_3 taken together form a six membered aromatic ring:

 R_7 and R_9 are independently selected from hydrogen, halogen, loweralkyl, haloloweralkyl, cycloalkyl, heterocyclyl, substituted heterocyclyl or heterocyclylalkyl; and R_1 , R_2 , R_3 , R_4 , R_7 , R_8 , R_9 , and R_{10} are absent when the ring atom to which they would otherwise be bonded is sulfur or double-bonded nitrogen: or

the a pharmaceutically acceptable salts, esters, or prodrugs thereof, provided that R_1 , R_2 , R_3 , R_4 , R_7 , R_8 , R_9 , and R_{10} are not all hydrogen when A, B, C, D, E, F, and H are carbon.

In one embodiment, the compounds of Formula (I) have a backbone structure wherein D is nitrogen, and A-C and E-H are carbon.

In one embodiment, when D is carbon, at least one, or at least two of R_1 - R_4 , and R_7 - R_{10} are not hydrogen.

In one embodiment, R_1 through R_4 , and R_8 and R_{10} are independently selected from at least two of the group consisting of hydrogen, halogen, loweralkyl, cycloalkyl, heterocyclyl, substituted heterocyclyl, alkylheterocyclyl, amino, imino, haloloweralkyl, alkoxy, nitro, alkylsulfonyl, arylalkyl, arylalkylaryl, arylaryl, aryloxy, arylamino, acylamino, acyloxyamino, alkylamino, alkylamino, alkylamino, alkylamino, alkylamino, alkylamino, alkylamino, alkylamino, alkylamino, alkoxyalkylamino, alkoxyalkylheterocyclyl, mercaptoalkoxyalkyl, cyano, formyl, -COOR11 where R_{11} is hydrogen, loweralkyl, aryl, heterocyclyl, monosaccharide or disaccharide,

and -CONR $_{12}$ R $_{13}$ where R $_{12}$ and R $_{13}$ are independently selected from hydrogen, loweralkyl, aryl, heterocyclyl, saccharide, peptide and amino acid residues; and R $_4$ is not present when D is nitrogen.

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In an additional embodiment, 4A, B, C, D, E, F, G, and H are independently selected from carbon and nitrogen;

 R_1 , R_2 , R_3 , R_4 , R_8 and R_{10} are independently selected from the group consisting of hydrogen, halogen, loweralkyl, alkyl, substituted alkyl, heterocyclyl, substituted heterocyclyl, substituted alkenyl, (substituted alkyl)(alkyl)amino, haloloweralkyl, hydroxy, alkoxy, substituted alkoxy, hydroxyalkylthio, nitro, N-alkylsulfonamide, cyano, -COOR $_{11}$ wherein R_{11} is hydrogen, loweralkyl, aryl, heterocyclyl, monosaccharide or disaccharide, and -CONR $_{12}R_{13}$ wherein R_{12} and R_{13} are independently selected from hydrogen, loweralkyl, aryl, heterocyclyl, saccharide, peptide and amino acid residues.

For the compounds described herein:

The term "loweralkyl" refers to branched or straight chain acyclical alkyl groups comprising one to ten carbon atoms, including, e.g., methyl, ethyl, propyl, isopropyl, n-butyl, t-butyl, neopentyl and the like.

The term "alkyl" refers to alkyl groups that do not contain heteroatoms. Thus the term includes straight chain alkyl groups such as methyl, ethyl, propyl, butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, undecyl, dodecyl and the like. The phrase also includes branched chain isomers of straight chain alkyl groups, including but not limited to, the following that are provided by way of example: -CH(CH₃)₂, -CH(CH₃)(CH₂CH₃), -CH(CH₂CH₃)₂, -C(CH₂CH₃)₃, -CH₂CH(CH₃)₂, -CH₂CH(CH₃)(CH₂CH₃), -CH₂CH(CH₃)₃, -CH₂C(CH₂CH₃)₃, -CH₂CH(CH₃)₃, -CH₂CH(CH₃)₃

 $CH_2CH_2CH_2CH_3)_2$, $-CH_2CH_2C(CH_3)_3$, $-CH_2CH_2C(CH_2CH_3)_3$, $-CH(CH_3)CH_2CH(CH_3)_2$, $-CH(CH_3)CH(CH_3)CH(CH_3)_2$.

CH(CH₂CH₃)CH(CH₃)(CH₂CH₃), and others. The term also includes cyclic alkyl groups such as cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclohetyl, and cyclooctyl and such rings substituted with straight and branched chain alkyl groups as defined above. The term also includes polycyclic alkyl groups such as, but not limited to, adamantyl norbornyl, and bicyclo[2.2.2]octyl and such rings substituted with straight and branched chain alkyl groups as defined above. Thus, the phrase unsubstituted alkyl groups includes primary alkyl groups, secondary alkyl groups, and tertiary alkyl groups. Unsubstituted alkyl groups may be bonded to one or more carbon atom(s), oxygen atom(s), nitrogen atom(s), and/or sulfur atom(s) in the parent compound. Preferred unsubstituted alkyl groups include straight and branched chain alkyl

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groups and cyclic alkyl groups having 1 to 20 carbon atoms. More preferred such unsubstituted alkyl groups have from 1 to 10 carbon atoms while even more preferred such groups have from 1 to 5 carbon atoms. Most preferred unsubstituted alkyl groups include straight and branched chain alkyl groups having from 1 to 3 carbon atoms and include methyl, ethyl, propyl, and — CH(CH₃)2.

The phrase "substituted alkyl" refers to an unsubstituted alkyl group as defined above in which one or more bonds to a carbon(s) or hydrogen(s) are replaced by a bond to non-hydrogen and non-carbon atoms such as, but not limited to, a halogen atom in halides such as F. Cl. Br. and I; a phosphorus atom in groups such as phosphate and dialkyl alkylphosphonate; oxygen atom in groups such as hydroxyl groups, alkoxy groups, aryloxy groups, and ester groups; a sulfur atom in groups such as thiol groups, alkyl and aryl sulfide groups, sulfone groups, sulfonyl groups, and sulfoxide groups; a nitrogen atom in groups such as amines, amides, alkylamines, dialkylamines, arylamines, alkylarylamines, diarylamines, N-oxides, imides, and enamines: a silicon atom in groups such as in trialkylsilyl groups, dialkylarylsilyl groups, alkyldiarylsilyl groups, and triarylsilyl groups; and other heteroatoms in various other groups. Substituted alkyl groups also include groups in which one or more bonds to a carbon(s) or hydrogen(s) atom is replaced by a bond to a heteroatom such as oxygen in carbonyl, carboxyl, and ester groups; nitrogen in groups such as imines, oximes, hydrazones, and nitriles. Preferred substituted alkyl groups include, among others, alkyl groups in which one or more bonds to a carbon or hydrogen atom is/are replaced by one or more bonds to fluorine atoms. One example of a substituted alkyl group is the trifluoromethyl group and other alkyl groups that contain the trifluoromethyl group. Other alkyl groups include those in which one or more bonds to a carbon or hydrogen atom is replaced by a bond to an oxygen atom such that the substituted alkyl group contains a hydroxyl, alkoxy, aryloxy group, or heterocyclyloxy group. Still other alkyl groups include alkyl groups that have an amine, alkylamine, dialkylamine, arylamine, (alkyl)(aryl)amine, diarylamine, heterocyclylamine, (alkyl)(heterocyclyl)amine, (aryl)(heterocyclyl)amine, or diheterocyclylamine group.

The term "alkoxy" refers to RO- wherein R, for example, is alkyl such as loweralkyl defined above. Representative examples of loweralkyl alkoxy groups include methoxy, ethoxy, t-butoxy and the like.

The phrase "substituted alkoxy" refers to RO-, where R is, for example, an alkyl substituted, for example, with a halogen. RO is for example OCF $_3$.

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The term "alkenyl" refers to a branched or straight chain groups comprising two to twenty carbon atoms that also comprises one or more carbon-carbon double bonds.

Representative alkenyl groups include prenyl, 2-propenyl (i.e., allyl), 3-methyl-2-butenyl, 3,7-dimethyl-2,6-octadienyl, 4,8-dimethyl-3,7-nonadienyl, 3,7,11-trimethyl-2,6,10-dodecatrienyl and the like.

The phrase "substituted alkenyl" refers to alkenyl groups that are substituted, for example, diethyl hex-5-enylphosponate, and others with an alkyl or substituted alkyl group such as dialkyl phosphate or an ester such as an acetate ester.

. The phrase "dialkyl amino" refers to an amino group substituted with two alkyl groups such as C1-20 alkyl groups.

The phrase "substituted dialkyl amino" refers to a dialkylamino substituted, for example, with a carboxylic acid, ester, hydroxy or alkoxy.

The term "hydroxyalkylthio" refers to a thio radical to which is appended a hydroxyalkyl group, where the alkyl is for example lower alkyl. An example is hydroxyethylthio, - SCH_2CH_2OH .

The term "N-alkylsulfonamide" refers to the group -SO $_2$ NHalkyl, where alkyl is, for example, octyl.

The term "alkynyl" refers to a branched or straight chain comprising two to twenty carbon atoms that also comprises one or more carbon-carbon triple bonds. Representative alkynyl groups include ethynyl, 2-propynyl (propargyl), 1-propynyl and the like.

The term "aryl" refers to aryl groups that do not contain heteroatoms. Thus the term includes, but is not limited to, groups such as phenyl, biphenyl, anthracenyl, naphthenyl by way of example. Although the phrase "unsubstituted aryl" includes groups containing condensed rings such as naphthalene, it does not include aryl groups that have other groups such as alkyl or halo groups bonded to one of the ring members, as aryl groups such as tolyl are considered herein to be substituted aryl groups as described below. A preferred unsubstituted aryl group is phenyl. Unsubstituted aryl groups may be bonded to one or more carbon atom(s), oxygen atom(s), nitrogen atom(s), and/or sulfur atom(s) in the parent compound, however.

The phrase "substituted aryl group" has the same meaning with respect to aryl groups that substituted alkyl groups had with respect to alkyl groups. However, a substituted aryl group also includes aryl groups in which one of the aromatic carbons is bonded to one of the non-carbon or non-hydrogen atoms described above and also includes aryl groups in which one or more aromatic carbons of the aryl group is bonded to a substituted and/or unsubstituted alkyl, alkenyl,

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or alkynyl group as defined herein. This includes bonding arrangements in which two carbon atoms of an aryl group are bonded to two atoms of an alkyl, alkenyl, or alkynyl group to define a fused ring system (e.g. dihydronaphthyl) or tetrahydronaphthyl). Thus, the phrase "substituted aryl" includes, but is not limited to tolyl, and hydroxyphenyl among others.

The term "arylalkyl" refers to a loweralkyl radical to which is appended an aryl group. Representative arylalkyl groups include benzyl, phenylethyl, hydroxybenzyl, fluorobenzyl, fluorophenylethyl and the like.

The phrase "unfused arylaryl" refers to a group or substituent to which two aryl groups. that are not condensed to each other, are bound. Exemplary unfused arylaryl compounds include. for example, phenylbenzene, diphenyldiazene, 4-methylthio-1-phenylbenzene, phenoxybenzene, (2-phenylethynyl)benzene, diphenyl ketone, (4-phenylbuta-1,3-diynyl)benzene. phenylbenzylamine. (phenylmethoxy)benzene, and the like. Preferred substituted unfused arvlaryl groups include: 2-(phenylamino)-N-[4-(2-phenylethynyl)phenyl]acetamide, 1.4-phenylethynyl)phenyl]propanamide, 2-amino-N-[4-(2-phenylethynyl)phenyl]acetamide, 2-(cyclopropylamino)-N-[4-(2-phenylethynyl)phenyl]acetamide, 2-(ethylamino)-N-[4-(2-phenylethynyl)phenyl]acetamide, 2-(ethylamino)-N-[4-(2-phenylethynyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethynyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethynyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylethyl)phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenyllacetamide, 2-(ethylamino)-N-[4-(2-phenylla phenylethynyl)phenyl]acetamide, 2-[(2-methylpropyl)amino]-N-[4-(2phenylethynyl)phenyl]acetamide, 5-phenyl-2H-benzo[d]1,3-dioxolene, 2-chloro-1-methoxy-4phenylbenzene, 2-[(imidazolylmethyl)amino]-N-[4-(2-phenylethynyl)phenyl]acetamide, 4phenyl-1-phenoxybenzene, N-(2-aminoethyl)[4-(2-phenylethynyl)phenyl]carboxamide, 2-{[(4fluorophenyl)methyl]amino}-N-[4-(2-phenylethynyl)phenyl]acetamide, 2-{[(4methylphenyl)methyl]amino}-N-[4-(2-phenylethynyl)phenyl]acetamide, 4-phenyl-1-(trifluoromethyl)benzene, 1-butyl-4-phenylbenzene, 2-(cyclohexylamino)-N-[4-(2phenylethynyl)phenyl]acetamide, 2-(ethylmethylamino)-N-[4-(2phenylethynyl)phenyl]acetamide, 2-(butylamino)-N-[4-(2-phenylethynyl)phenyl]acetamide, N-[4-(2-phenylethynyl)phenyl]-2-(4-pyridylamino)acetamide, N-[4-(2-phenylethynyl)phenyl]-2-(quinuclidin-3-ylamino)acetamide, N-[4-(2-phenylethynyl)phenyl]pyrrolidin-2-ylcarboxamide, 2-amino-3-methyl-N-[4-(2-phenylethynyl)phenyl]butanamide, 4-(4-phenylbuta-1,3divnyl)phenylamine, 2-(dimethylamino)-N-[4-(4-phenylbuta-1,3-diynyl)phenyl]acetamide, 2-(ethylamino)-N-[4-(4-phenylbuta-1,3-diynyl)phenyl]acetamide, 4-ethyl-1-phenylbenzene, 1-[4-(2-phenylethynyl)phenyl]ethan-1-one, N-(1-carbamoyl-2-hydroxypropyl)[4-(4-phenylbuta-1.3divnyl)phenyl]carboxamide, N-[4-(2-phenylethynyl)phenyl]propanamide, 4-methoxyphenyl

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(3-phenylphenoxy)ethanehydroxamic acid, 3-phenylphenyl propanoate, 1-(4-ethoxyphenyl)-4-methoxybenzene, and [4-(2-phenylethynyl)phenyl]pyrrole.

The phrase "unfused heteroarylaryl" refers to a unfused arylaryl group where one of the aryl groups is a heteroaryl group. Exemplary heteroarylaryl groups include, for example, 2-phenylpyridine, phenylpyrrole, 3-(2-phenylethynyl)pyridine, phenylpyrazole, 5-(2-phenylethynyl)-1,3-dihydropyrimidine-2,4-dione, 4-phenyl-1,2,3-thiadiazole, 2-(2-phenylethynyl)pyrazine, 2-phenylthiophene, phenylimidazole, 3-(2-piperazinylphenyl)furan, 3-(2,4-dichlorophenyl)-4-methylpyrrole, and the like. Preferred substituted unfused heteroarylaryl groups include: 5-(2-phenylethynyl)pyrimidine-2-ylamine, 1-methoxy-4-(2-thienyl)benzene, 1-methoxy-3-(2-thienyl)benzene, 5-methyl-2-phenylpyridine, 5-methyl-3-phenylisoxazole, 2-[3-(trifluoromethyl)phenyl]furan, 3-fluoro-5-(2-furyl)-2-methoxy-1-prop-2-enylbenzene, (hydroxyimino)(5-phenyl(2-thienyl))methane, 5-[(4-methylpiperazinyl)methyl]-2-phenylthiophene, 2-(4-ethylphenyl)thiophene, 4-methylthio-1-(2-thienyl)benzene, 2-(3-nitrophenyl)thiophene, (tert-butoxy)-N-[(5-phenyl(3-pyridyl))methyl]carboxamide, hydroxy-N-[(5-phenyl(3-pyridyl))methyl]amide, 2-(phenylmethylthio)pyridine, and benzylimidazole.

The phrase "unfused heteroarylheteroaryl" refers to an unfused arylaryl group where both of the aryl groups is a heteroaryl group. Exemplary heteroarylheteroaryl groups include, for example, 3-pyridylimidazole, 2-imidazolylpyrazine, and the like. Preferred substituted unfused heteroarylheteroaryl groups include: 2-(4-piperazinyl-3-pyridyl)furan, diethyl(3-pyrazin-2-yl(4-pyridyl))amine, and dimethyl{2-[2-(5-methylpyrazin-2-yl)ethynyl](4-pyridyl)}amine.

The phrase "fused arylaryl" refers to an aryl group as previously defined that is condensed, and fully conjugated to an aryl group. Representative fused arylaryl groups include biphenyl, 4-(1-naphthyl)phenyl, 4-(2-naphthyl)phenyl and the like.

The phrase "fused heteroarylaryl" refers to an aryl group as previously defined that is condensed, and fully conjugated with a heteroaryl group. Representative fused heteroarylaryl groups include quinoline, quinazoline and the like.

The phrase "fused heteroarylheteroaryl" refers to a heteroaryl group as previously defined that is condensed, and fully conjugated with another heteroaryl group. Representative fused heteroarylheteroaryl groups include pyrazalopyrimidine, imidazoquinoline and the like.

The term "aryloxy" refers to RO- wherein R is an aryl group. Representative arylalkoxy group include benzyloxy, phenylethoxy and the like.

The term "arylalkoxy" refers to a lower alkoxy radical to which is appended an aryl group. Representative arylalkoxy group include benzyloxy, phenylethoxy and the like.

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The term "aryloxyaryl" refers to an aryl radical to which is appended an aryloxy group. Representative aryloxyaryl groups include 4-phenoxyphenyl, 3-phenoxyphenyl, 4-phenoxy-1-naphthyl, 3-phenoxy-1-naphthyl and the like.

The term "aryloxyarylalkyl" refers to an arylalkyl radical to which is appended an aryloxy group. Representative aryloxyarylalkyl groups include 4-phenoxyphenylmethyl, 3-phenoxyphenylmethyl, 4-phenoxyphenylethyl, 3-phenoxy-phenylethyl and the like.

The term "arylalkoxyaryl" refers to an aryl radical to which is appended an arylalkoxy group. Representative arylalkoxyaryl groups include 4-benzyloxylphenyl, 3-benzyloxyphenyl and the like.

The term "arylalkoxyarylalkyl" refers to an arylalkyl radical to which is appended an arylalkoxy group. Representative arylalkoxyarylalkyl groups include 4-benzyloxylbenzyl, 3-benzyloxybenzyl and the like.

The term "cycloalkyl" refers to an alicyclic group comprising from 3 to 7 carbon atoms including, but not limited to, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl and the like.

The term "cycloalkylalkyl" refers to a loweralkyl radical to which is appended a cycloalkyl group. Representative examples of cycloalkylalkyl include cyclopropylmethyl, cyclohexylmethyl, 2-(cyclopropyl)ethyl and the like.

The term "halogen" refers to iodine, bromine, chlorine or fluorine; "halo" refers to iodo, bromo, chloro or fluoro.

The term "haloalkyl" refers to a lower alkyl radical, as defined above, bearing at least one halogen substituent, for example, chloromethyl, fluoroethyl or trifluoromethyl and the like.

The term "heterocyclyl" (or heterocyclic, or heterocyclo) refers to both aromatic and nonaromatic ring compounds including monocyclic, bicyclic, and polycyclic ring compounds such as, but not limited to, quinuclidyl, containing 3 or more ring members of which one or more is a heteroatom such as, but not limited to, N, O, and S. Although the phrase "unsubstituted heterocyclyl" includes condensed heterocyclic rings such as benzimidazolyl, it does not include heterocyclyl groups that have other groups such as alkyl or halo groups bonded to one of the ring members as compounds such as 2-methylbenzimidazolyl are substituted heterocyclyl groups. Examples of heterocyclyl groups include, but are not limited to: unsaturated 3 to 8 membered rings containing 1 to 4 nitrogen atoms such as, but not limited to pyrrolyl, pyrrolinyl, imidazolyl, pyrazolyl, pyridyl, dihydropyridyl, pyrimidyl, pyrazinyl, pyridazinyl, triazolyl (e.g. 4H-1,2,4-triazolyl, 1H-1,2,3-triazolyl, 2H-1,2,3-triazolyl etc.), tetrazolyl, (e.g. 1H-tetrazolyl, 2H tetrazolyl, etc.); saturated 3 to 8 membered rings containing 1 to 4 nitrogen atoms such as, but

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not limited to, pyrrolidinyl, imidazolidinyl, piperidinyl, piperazinyl; condensed unsaturated heterocyclic groups containing 1 to 4 nitrogen atoms such as, but not limited to, indolyl, isoindolyl, indolinyl, indolizinyl, benzimidazolyl, quinolyl, isoquinolyl, indazolyl, benzotriazolyl; unsaturated 3 to 8 membered rings containing 1 to 2 oxygen atoms and 1 to 3 nitrogen atoms such as, but not limited to, oxazolyl, isoxazolyl, oxadiazolyl (e.g. 1,2.4oxadiazolyl, 1,3,4-oxadiazolyl, 1,2,5-oxadiazolyl, etc.); saturated 3 to 8 membered rings containing 1 to 2 oxygen atoms and 1 to 3 nitrogen atoms such as, but not limited to, morpholinyl; unsaturated condensed heterocyclic groups containing 1 to 2 oxygen atoms and 1 to 3 nitrogen atoms, for example, benzoxazolyl, benzoxadiazolyl, benzoxazinyl (e.g. 2H-1.4benzoxazinyl etc.); unsaturated 3 to 8 membered rings containing 1 to 3 sulfur atoms and 1 to 3 nitrogen atoms such as, but not limited to, thiazolyl, isothiazolyl, thiadiazolyl (e.g. 1,2.3thiadiazolyl, 1,2,4-thiadiazolyl, 1,3,4-thiadiazolyl, 1,2,5-thiadiazolyl, etc.); saturated 3 to 8 membered rings containing 1 to 2 sulfur atoms and 1 to 3 nitrogen atoms such as, but not limited to, thiazolodinyl; saturated and unsaturated 3 to 8 membered rings containing 1 to 2 sulfur atoms such as, but not limited to, thienyl, dihydrodithiinyl, dihydrodithionyl, tetrahydrothiophene, tetrahydrothiopyran; unsaturated condensed heterocyclic rings containing 1 to 2 sulfur atoms and 1 to 3 nitrogen atoms such as, but not limited to, benzothiazolyl, benzothiadiazolyl, benzothiazinyl (e.g. 2H-1,4-benzothiazinyl, etc.), dihydrobenzothiazinyl (e.g. 2H-3,4dihydrobenzothiazinyl, etc.), unsaturated 3 to 8 membered rings containing oxygen atoms such as, but not limited to furyl; unsaturated condensed heterocyclic rings containing 1 to 2 oxygen atoms such as benzodioxolyl (e.g. 1,3-benzodioxoyl, etc.); unsaturated 3 to 8 membered rings containing an oxygen atom and 1 to 2 sulfur atoms such as, but not limited to, dihydrooxathiinyl; saturated 3 to 8 membered rings containing 1 to 2 oxygen atoms and 1 to 2 sulfur atoms such as 1,4-oxathiane; unsaturated condensed rings containing 1 to 2 sulfur atoms such as benzothienyl, benzodithiinyl; and unsaturated condensed heterocyclic rings containing an oxygen atom and 1 to 2 oxygen atoms such as benzoxathiinyl. Heterocyclyl group also include those described above in which one or more S atoms in the ring is double-bonded to one or two oxygen atoms (sulfoxides and sulfones). For example, heterocyclyl groups include tetrahydrothiophene, tetrahydrothiophene oxide, and tetrahydrothiophene 1,1-dioxide. Preferred heterocyclyl groups contain 5 or 6 ring members. More preferred heterocyclyl groups include morpholine. piperazine, piperidine, pyrrolidine, imidazole, pyrazole, 1,2,3-triazole, 1,2,4-triazole, tetrazole, thiomorpholine, thiomorpholine in which the S atom of the thiomorpholine is bonded to one or

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more O atoms, pyrrole, homopiperazine, oxazolidin-2-one, pyrrolidin-2-one, oxazole, quinuclidine, thiazole, isoxazole, furan, and tetrahydrofuran.

The phrase "substituted heterocyclyl" refers to an heterocyclyl group as defined above in which one of the ring members is bonded to a non-hydrogen atom such as described above with respect to substituted alkyl groups and substituted aryl groups. Examples, include, but are not limited to, 2-methylbenzimidazolyl, 5-methylbenzimidazolyl, 5-chlorobenzthiazolyl, 1-methyl piperazinyl, and 2-chloropyridyl among others.

"Aminosulfonyl" refers to the group $-S(O)_2$ -NH₂. "Substituted aminosulfonyl" refers to the group $-S(O)_2$ -NRR' where R is loweralkyl and R' is hydrogen or a loweralkyl. The term "aralkylaminosulfonlyaryl" refers to the group -aryl- $S(O)_2$ -NH-aralkyl, where the aralkyl is loweraralkyl.

"Carbonyl" refers to the divalent group -C(O)-.

"Carbonyloxy" refers generally to the group -C(O)-O-,. Such groups include esters, -C(O)-O-R, where R is loweralkyl, cycloalkyl, aryl, or loweraralkyl. The term "carbonyloxycycloalkyl" refers generally to both an "carbonyloxycarbocycloalkyl" and an "carbonyloxyheterocycloalkyl", i.e., where R is a carbocycloalkyl or heterocycloalkyl, respectively. The term "arylcarbonyloxy" refers to the group -C(O)-O-aryl, where aryl is a mono- or polycyclic, carbocycloaryl or heterocycloaryl. The term "aralkylcarbonyloxy" refers to the group -C(O)-O-aralkyl, where the aralkyl is loweraralkyl.

The term "sulfonyl" refers to the group $-SO_2$. "Alkylsulfonyl" refers to a substituted sulfonyl of the structure $-SO_2R$ - in which R is alkyl. Alkylsulfonyl groups employed in compounds of the present invention are typically loweralkylsulfonyl groups having from 1 to 6 carbon atoms in its backbone structure. Thus, typical alkylsulfonyl groups employed in compounds of the present invention include, for example, methylsulfonyl (i.e., where R is methyl), ethylsulfonyl (i.e., where R is ethyl), propylsulfonyl (i.e., where R is propyl), and the like. The term "arylsulfonyl" refers to the group $-SO_2$ -aryl. The term "aralkylsulfonyl" refers to the group $-SO_2$ -aralkyl, in which the aralkyl is loweraralkyl. The term "sulfonamido" refers to $-SO_2$ NH2.

The term "carbonylamino" refers to the divalent group -NH-C(O)- in which the hydrogen atom of the amide nitrogen of the carbonylamino group can be replaced a loweralkyl, aryl, or loweraralkyl group. Such groups include moieties such as carbamate esters (-NH-C(O)-O-R) and amides -NH-C(O)-O-R, where R is a straight or branched chain loweralkyl, cycloalkyl, or aryl or loweraralkyl. The term "loweralkylcarbonylamino" refers to alkylcarbonylamino where R is a

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loweralkyl having from 1 to about 6 carbon atoms in its backbone structure. The term "arylcarbonylamino" refers to group –NH-C(O)-R where R is an aryl. Similarly, the term "aralkylcarbonylamino" refers to carbonylamino where R is a lower aralkyl.

The term "guanidino" or "guanidyl" refers to moieties derived from guanidine,

H2N-C(=NH)-NH₂. Such moieties include those bonded at the nitrogen atom carrying the formal
double bond (the "2"-position of the guanidine, e.g., diaminomethyleneamino, (H2N)₂C=NH-)
and those bonded at either of the nitrogen atoms carrying a formal single bond (the "1-" and/or
"3"-positions of the guandine, e.g., H₂N-C(=NH)-NH-). The hydrogen atoms at any of the
nitrogens can be replaced with a suitable substituent, such as loweralkyl, aryl, or loweraralkyl.

Representative cycloimido and heterocycloimido groups include, for example, those shown below. These cycloimido and heterocycloimido can be further substituted and may be attached at various positions as will be apparent to those having skill in the organic and medicinal chemistry arts in conjunction with the disclosure herein.

Representative substituted amidino and heterocycloamidino groups include, for example, those shown below. These amidino and heterocycloamidino groups can be further substituted as will be apparent to those having skill in the organic and medicinal chemistry arts in conjunction with the disclosure herein.

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Representative substituted alkylcarbonylamino, alkyloxycarbonylamino, aminoalkyloxycarbonylamino, and arylcarbonylamino groups include, for example, those shown below. These groups can be further substituted as will be apparent to those having skill in the organic and medicinal chemistry arts in conjunction with the disclosure herein.

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Representative substituted aminocarbonyl groups include, for example, those shown below. These can heterocyclo groups be further substituted as will be apparent to those having skill in the organic and medicinal chemistry arts in conjunction with the disclosure herein.

Representative substituted alkoxycarbonyl groups include, for example, those shown below. These alkoxycarbonyl groups can be further substituted as will be apparent to those having skill in the organic and medicinal chemistry arts in conjunction with the disclosure herein.

"Substituted" refers to the definite replacement of hydrogen with one or more monovalent or divalent radicals. Suitable substitution groups include, those described herein for particular groups, as well as hydroxyl, nitro, amino, imino, cyano, halo, thio, thioamido, amidino, imidino, oxo, oxamidino, methoxamidino, imidino, guanidino, sulfonamido, carboxyl, formyl, alkyl, substituted alkyl, haloloweralkyl, loweralkoxy, haloloweralkoxy, loweralkoxy, alkyl, alkylcarbonyl, arylcarbonyl, aralkylcarbonyl, heteroarylcarbonyl, heteroaralkylcarbonyl, alkylthio, aminoalkyl, cyanoalkyl, benzyl, pyridyl, pyrazolyl, pyrrole, thiophene, imidazolyl, and the like.

The term "linking moiety" refers to a covalent bond or an uncyclized divalent group, such as, for example, -CO-, -O-, -S-, -CH₂-, -NH-, and substituted or unsubstituted alkyl, alkenyl, alkynyl, carbonyl, alkoxycarbonyl groups as defined herein.

The term "SMIP compound" refers to small molecule immunopotentiating compounds, that include small molecule compounds below about MW 1000 g/mol, preferably MW 800 g/mol that are capable of stimulating or modulating a pro-inflammatory response in a patient. In an embodiment, the SMIP compounds are able to stimulate human peripheral blood mononuclear cells to produce cytokines. Preferred SMIP compounds and derivatives thereof include, for example, aminoazavinyl compounds, benzazole compounds, acylpiperazine compounds, indoledione compounds, tetrahydroisoquinoline (THIQ) compounds, anthraquinone compounds, indanedione compounds, pthalimide compounds, benzocyclodione compounds, aminobenzimidazole quinolinone (ABIQ) compounds, hydraphthalimide compounds, pyrazolopyrimidine compounds, quinazilinone compounds, quinoxaline compounds, tetrahydropyrrolidinoquinoxaline compounds, pyrrole compounds, benzophenone compounds, sterol compound, and isoxazole compounds.

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The term "SMIS compound" refers to small molecule immunosuppressant compounds, that include small molecule compounds below about about MW 1000 g/mol, preferably MW 800 g/mol, capable of suppressing or modulating a pro-inflammatory response in a patient.

Acylpiperazine compounds as described throughout this application include compounds of formula (III) as shown below:

$$\begin{array}{c} D_{3} \\ D_{2} \\ N \\ N \\ N \\ (R_{10})_{n} \\ \end{array}$$

wherein.

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 R_9 is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, arylalkyl, arylalkenyl, heteroarylalkyl, and heteroarylalkenyl;

R₁₀ is substituted or unsubstituted alkyl;

n is an integer from 0-2; and

if D_1 is carbon than D_2 is oxygen, D_3 is absent, and D_4 is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, carbocycyl, alkoxyaryl, fused arylaryl, fused arylheteroaryl, and fused heteroarylaryl; or,

if D_1 is nitrogen than D_2 is nitrogen, D_4 is absent, and D_3 is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, carbocycyl, alkoxyaryl, fused arylaryl, fused arylaryl, fused arylaryl, and fused heteroarylaryl.

. Indoledione compounds as described throughout this application include compounds of formula (IV) as shown below:

5 wherein.

 R_{11} and R_{12} are independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylakoxy, alkylamino, arylalkylamino, arylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups; and,

 R_{13} is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, arylalkyl, heteroarylalkyl, heterocyclyl, heterocyclylalkyl, and alkylbenzyl.

Tetrahydroisoquinoline (THIQ) compounds as described throughout this application include compounds of formula (V) as shown below:

wherein.

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L is a covalent bond or selected from the group consisting of -CH₂-, -CO-, -O-, -S-, CHF, -NH-, -NR₂₀-, where R₂₀ is lower alkyl;

 R_{14} is selected from the group consisting of hydrogen, halogen, and substituted or unsubstituted alkyl;

 R_{15} is selected from the group consisting of substituted or unsubstituted carbocyclyl, aryl, arylalkyl, alkoxyaryl, heteroaryl, heterocyclyl;

 R_{16} is selected from the group consisting of hydrogen, halogen, and substituted or unsubstituted alkyl;

 R_{17} is selected from the group consisting of hydrogen, halogen, and substituted or unsubstituted alkyl; and,

R₁₈ and R₁₉ are independently selected from the group consisting of H, hydroxy, halogen, alkoxy, amino, unsubstituted alkyl, substituted alkyl, and alkylamino.

Benzocyclodione compounds as described throughout this application include

compounds of formula (VI) as shown below:

wherein.

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E is selected from the group consisting of NR₂₅ or CR₂₆R₂₇;

R₂₁, R₂₃, and R₂₄ are independently selected from the group consisting of H, hydroxy, halogen, alkoxy, amino, unsubstituted alkyl, substituted alkyl, and alkylamino;
R₂₂ is selected from the group consisting or H, hydroxy, halogen, alkoxy, amino, and unsubstituted or substituted alkyl, and alkylamino, arylalkyl, heteroarylalkyl, aryl, heteroaryl, arylcarbonyl, heterocyclyl, heterocyclylalkyl, and heteroarylcarbonyl;
R₂₅ is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, heterocyclyl, carbocyclyl, arylalkyl, heteroarylalkyl, and heterocyclyalkyl;
R₂₆ is selected from the group consisting of H, halogen, hydroxy, amino, and substituted or unsubstituted alkyl, carbonylalkyl, and alkylcarbonylalkyl; and,
R₂₇ is selected from the group aryl, arylalkyl, heteroarylalkyl, heterocyclyl, heterocyclylalkyl, carbocyclyl, arylcarbonylalkyl, and arylalkylcarbonyl.

Aminoazavinyl compounds as described throughout this application include compounds of formula (VII) as shown below:

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wherein,

G is either S or NH;

 R_{28} is selected from the group consisting of H, and substituted or unsubstituted alkyl, aryl, heteroaryl, heteroarylalkyl, arylalkyl, carbocyclyl, carbocyclylalkyl, heterocyclyl, and heterocyclylalkyl;

Q is selected from the group consisting of hydrogen, substituted alkyl, unsubstituted alkyl, and aryl, substituted aryl, heteroaryl, substituted heteroaryl, heterocyclyl, substituted

heterocyclyl, fused or unfused arylaryl, substituted arylaryl, arylheteroaryl, substituted arylheteroaryl, heteroarylheteroaryl, and substituted heteroarylheteroaryl:

 V_1 is selected from the group consisting of alkyl, substituted alkyl, aryl, substituted aryl, arylalkyl, substituted arylalkyl, heteroarylalkyl, substituted heteroarylalkyl, substituted heteroarylalkyl, substituted heteroarylalkyl, substituted heteroarylalkyl, substituted heteroarylalkyl, alkoxy, substituted alkoxy, aminocarbonyl, alkoxycarbonyl, carboxyl sulfonyl, methanesulfonyl, and substituted or unsubstituted alkylcarbonyl, arylcarbonyl, arylcarbonyl, heteroarylcarbonyl, heteroaralkylcarbonyl, arylcarbonyloxy, aralkylcarbonyloxy, heteroarylcarbonyloxy, heteroarylcarbonyloxy, formyl, loweralkylcarbonyloxy, alkylaminocarbonyloxy, arylaminocarbonyl, aminoaryl, alkylsulfonyl, sulfonamido, aminoalkoxy, alkylamino, heteroarylamino, alkylcarbonylamino, alkylaminocarbonylamino, arylaminocarbonylamino, aralkylcarbonylamino, heteroarylcarbonylamino, arylaminocarbonylamino, cycloamidino, cycloalkyl, cycloimido, arylsulfonyl and arylsulfonamido; and, V_2 is selected from the group consisting of hydrodgen, halogen, alkyl, substituted alkyl,

 V_2 is selected from the group consisting of hydrodgen, halogen, alkyl, substituted alkyl, aryl, substituted aryl, arylalkyl, substituted arylalkyl, heteroaryl, substituted heteroaryl, heteroarylalkyl, substituted heteroarylalkyl, substituted alkoxy, aminocarbonyl, alkoxycarbonyl, carboxyl sulfonyl, methanesulfonyl, and substituted or unsubstituted alkylcarbonyl, arylcarbonyl, aralkylcarbonyl, heteroarylcarbonyl, heteroaralkylcarbonyl, alkylcarbonyloxy, aralkylcarbonyloxy, heteroarylcarbonyloxy, heteroaralkylcarbonyloxy, alkylaminocarbonyloxy, arylaminocarbonyloxy, formyl, loweralkylcarbonyl, loweralkoxycarbonyl, aminoarbonyl, aminoaryl, alkylsulfonyl, sulfonamido, aminoalkoxy, alkylamino, heteroarylamino, alkylcarbonylamino, alkylamino, arylaminocarbonylamino, arylaminocarbonylamino, aralkylcarbonylamino, heteroarylcarbonylamino, arylcarbonylamino, cycloamidino, cycloalkyl, cycloimido, arylsulfonyl and arylsulfonamido.

Lactam compounds as described throughout this application include compounds of formula (VIII) as shown below:

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wherein.

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W₁ is selected from the group consisting of -OH, -OR₃₆ groups, -NR₃₇R₃₈;
W₂ is selected from the group consisting of O, S, and NR₃₉ groups;

 R_{29} and R_{30} join to form a 5 to 6 membered substituted or unsubstituted ring comprising all carbon atoms or at least one O, N, or S atom;

 R_{35} and R_{39} may be the same or different and are selected from the group consisting of H, -OH substituted and unsubstituted alkyl groups, substituted and unsubstituted aryl groups, -C(=O)H, -C(=O)-alkyl groups, and -C(=O)-aryl groups;

 R_{31} , R_{32} , R_{33} , and R_{34} may be the same or different and are independently selected from the group consisting of H, Cl, Br, F, I, -NO₂, -CN, -OH, -OR₄₀ groups, -NR₄₁R₄₂ groups, -C(=O)R₄₃ groups, -SH groups, substituted and unsubstituted amidinyl groups, substituted and unsubstituted alkyl groups, substituted and unsubstituted alkenyl groups, substituted and unsubstituted arylaminoalkyl groups, substituted and unsubstituted arylaminoalkyl groups, substituted and unsubstituted and unsubstituted and unsubstituted and unsubstituted (alkyl)(aryl)aminoalkyl groups, substituted aminoalkyl groups, substituted and unsubstituted diheterocyclylaminoalkyl groups, substituted and unsubstituted (alkyl)(heterocyclylaminoalkyl groups, substituted and unsubstituted (alkyl)(heterocyclyl)aminoalkyl groups, substituted and unsubstituted (aryl)(heterocyclyl)aminoalkyl groups, substituted and unsubstituted (aryl)(heterocyclyl)aminoalkyl groups, substituted and unsubstituted (aryl)(heterocyclyl)aminoalkyl groups, substituted and unsubstituted hydroxyalkyl

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groups, substituted and unsubstituted alkoxyalkyl groups, substituted and unsubstituted aryloxyalkyl groups, and substituted and unsubstituted heterocyclyloxyalkyl groups; R_{36} is selected from the group consisting of substituted and unsubstituted alkyl groups, substituted and unsubstituted aryl groups, substituted and unsubstituted heterocyclyl

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groups, substituted and unsubstituted heterocyclylalkyl groups, -C(=O)H, -C(=O)-alkyl groups, -C(=O)-aryl groups, -C(=O)-alkyl groups, -C(=O)-aryl groups, $-C(=O)NH_2$, -C(=O)NH(alkyl) groups, -C(=O)NH(aryl) groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-NH_2$, $-NH(alkyl)_2$ groups, $-C(=O)NH(alkyl)_2$ groups, $-C(=O)NH(alkyl)_2$ groups, $-C(=O)NH(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, $-C(=O)N(alkyl)_2$ groups, and $-C(=O)N(alkyl)_2$ groups, substituted and unsubstituted alkyl groups, substituted and unsubstituted aryl groups, and substituted and unsubstituted heterocyclyl groups;

R₃₈ is selected from the group consisting of H, substituted and unsubstituted alkyl groups. substituted and unsubstituted aryl groups, substituted and unsubstituted heterocyclyl groups, -OH, alkoxy groups, aryloxy groups, -NH2, substituted and unsubstituted heterocyclylalkyl groups, substituted and unsubstituted aminoalkyl groups, substituted and unsubstituted alkylaminoalkyl groups, substituted and unsubstituted dialkylaminoalkyl groups, substituted and unsubstituted arylaminoalkyl groups. substituted and unsubstituted diarylaminoalkyl groups, substituted and unsubstituted (alkyl)(aryl)aminoalkyl groups, substituted and unsubstituted alkylamino groups, substituted and unsubstituted arylamino groups, substituted and unsubstituted dialkylamino groups, substituted and unsubstituted diarylamino groups, substituted and unsubstituted (alkyl)(aryl)amino groups, -C(=O)H, -C(=O)-alkyl groups, -C(=O)-aryl groups, -C(=O)O-alkyl groups, -C(=O)O-aryl groups, -C(=O)NH2, -C(=O)NH(alkyl) groups, -C(=O)NH(aryl) groups, -C(=O)N(alkyl)2 groups, -C(=O)N(aryl)2 groups, -C(=O)N(alkyl)(aryl) groups, -C(=O)-heterocyclyl groups, -C(=O)-Oheterocyclyl groups, -C(=O)NH(heterocyclyl) groups, -C(=O)-N(heterocyclyl)2 groups, -C(=O)-N(alkyl)(heterocyclyl) groups, -C(=O)-N(aryl)(heterocyclyl) groups, substituted and unsubstituted heterocyclylaminoalkyl groups, substituted and unsubstituted diheterocyclylaminoalkyl groups, substituted and unsubstituted (alkyl)(heterocyclyl)aminoalkyl groups, substituted and unsubstituted (aryl)(heterocyclyl)aminoalkyl groups, substituted and unsubstituted hydroxyalkyl groups, substituted and unsubstituted alkoxyalkyl groups, substituted and unsubstituted

aryloxyalkyl groups, and substituted and unsubstituted heterocyclyloxyalkyl groups;

 R_{41} is selected from the group consisting of H, substituted and unsubstituted alkyl groups, substituted and unsubstituted aryl groups, and substituted and unsubstituted heterocyclyl groups;

R₄₂ is selected from the group consisting of H, substituted and unsubstituted alkyl groups. substituted and unsubstituted aryl groups, substituted and unsubstituted heterocyclyl groups. -C(=O)H, -C(=O)-alkyl groups, -C(=O)-aryl groups, -C(=O)NH₂, -C(=O)NH(alkyl) groups, -C(=O)NH(aryl) groups, -C(=O)N(alkyl)₂ groups. -C(=O)N(aryl)₂ groups, -C(=O)N(alkyl)(aryl) groups, -C(=O)O-alkyl groups, -C(=O)O-aryl groups, substituted and unsubstituted aminoalkyl groups. substituted and unsubstituted alkylaminoalkyl groups, substituted and unsubstituted dialkylaminoalkyl groups, substituted and unsubstituted arylaminoalkyl groups, substituted and unsubstituted diarylaminoalkyl groups, substituted and unsubstituted (alkyl)(aryl)aminoalkyl groups, substituted and unsubstituted heterocyclylalkyl groups, -C(=O)-heterocyclyl groups, -C(=O)-O-heterocyclyl groups, -C(=O)NH(heterocyclyl) groups, -C(=O)-N(heterocyclyl)2 groups, -C(=O)-N(alkyl)(heterocyclyl) groups, -C(=O)-N(aryl)(heterocyclyl) groups, substituted and unsubstituted heterocyclylaminoalkyl groups, substituted and unsubstituted diheterocyclylaminoalkyl groups, substituted and unsubstituted (heterocyclyl)(alkyl)aminoalkyl groups, substituted and unsubstituted (heterocyclyl)(aryl)aminoalkyl groups, substituted and unsubstituted hydroxyalkyl groups, substituted and unsubstituted alkoxyalkyl groups, substituted and unsubstituted aryloxyalkyl groups, and substituted and unsubstituted heterocyclyloxyalkyl groups; and R₄₂ is selected from the group consisting of H, -NH₂, -NH(alkyl) groups, -NH(aryl) groups, -N(alkyl)2 groups, -N(aryl)2 groups, -N(alkyl)(aryl) groups, -NH(heterocyclyl) groups, -N(heterocyclyl)(alkyl) groups, -N(heterocyclyl)(aryl) groups, -N(heterocyclyl)2 groups, substituted and unsubstituted alkyl groups, substituted and unsubstituted arvi groups, -OH, substituted and unsubstituted alkoxy groups, substituted and unsubstituted heterocyclyl groups, substituted and unsubstituted aryloxy groups, heterocyclyloxy groups, -NHOH, -N(alkyl)OH groups, -N(aryl)OH groups, -N(alkyl)O-alkyl groups, -N(aryl)O-alkyl groups, -N(alkyl)O-aryl groups, and -N(aryl)O-aryl groups.

Preferably R₂₉ and R₃₀ join together to form a substituted or unsubstituted phenyl ring.

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Hydropthalamide compounds as described throughout this application include compounds of formula (IX) as shown below:

5 wherein,

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R₄₄ is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, arylalkyl, heteroarylalkyl, fused arylaryl, unfused arylaryl, fused heteroarylaryl, unfused heteroarylaryl, fused arylheteroaryl;

R45, R47, R49, and R51 may be the same or different and are independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylamino, arylalkoxy, alkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl; and R46, R48, R50, and R52 may be the same or different and are independently selected from the group consisting of H, halogen, and substituted or unsubstituted alkyl groups.

Benzophenone compounds as described throughout this application include compounds of formula (X) as shown below:

wherein.

R₅₃ is independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylalkylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl;

R₅₄ is independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl; and o and p are integers from 0-4.

Isoxazole compounds as described throughout this application include compounds of formula (XI) as shown below:

wherein.

R₅₅ is selected from the group consisting of substituted or unsubstituted aryl, arylalkyl, heteroaryl, heteroarylalkyl, heterocyclyl, and heterocyclylalkyl;

 R_{56} is selected from the group consisting of substituted or unsubstituted aryl, arylalkyl, heteroaryl, heteroarylalkyl, heterocyclyl, and heterocyclylalkyl; and, R_{57} is selected from the group consisting of H, halogen, hydoxy, and substituted or

unsubstituted alkyl, aryl, heteroaryl, heterocyclyl, and carbonyl.

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Sterol compounds as described throughout this application include compounds of formula (XII) as shown below:

5 wherein.

 R_{58} is selected from the group consisting of substituted or unsubstituted aryl, arylalkyl, heteroaryl, heteroarylalkyl, heterocyclyl, and heterocyclylalkyl. Preferably R_{58} is a pyranone substituent.

Quinazilinone compounds as described throughout this application include compounds of formula (XIII) as shown below:

5 wherein,

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 R_{59} is selected from the group consisting of H, halogen, hydroxy, and substituted or unsubstituted alkyl, aminoalkyl, alklyaminoalkyl, alkoxy, dialkylaminoalkyl, hydroxyalkyl, alkenyl, alkynyl, carbocyclyl, carbocyclylalkyl, heterocyclyl, and heterocyclylalkyl;

 R_{60} is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, arylalkyl, heteroarylalkyl, and heterocyclylalkyl; and,

 R_{61} , R_{62} , R_{63} , and R_{64} may be the same or different and are independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl,

alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups.

Pyrrole compounds as described throughout this application include compounds of formula (XIV) as shown below:

wherein,

 R_{65} is selected from the group consisting of H, hydroxy, and substituted or unsubstituted alkyl, aryl, heteroaryla, heteroarylalkyl, arylalkyl, heteroarylaminoalkyl, arylaminoalkyl, heteroaryloxyalkyl, and aryloxyalkyl groups;

 R_{66} , R_{67} , R_{68} , and R_{69} may be the same or different and are independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and -247-

substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylamino, alkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkoxy, heterocyclylalkoxy, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups.

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Further preferred pyrrole compounds include those shown in Formula (XV):

$$Ar \xrightarrow{R_{70}} \xrightarrow{R_{73}} \xrightarrow{R_{75}} \xrightarrow{R_{76}} \xrightarrow$$

lO

wherein:

K₁ is nitrogen, oxygen, or optionally substituted carbon:

W is absent or is selected from the group consisting of -O-, -S-, -S(O)-, -SO2-, -NH-, -NH-

.5 CO-, -NR'CO-, -NHSO₂-, -NR'SO₂-, -CO-, -CO₂--, -CH₂--, -CF₂--, CHF, -CONH-, -CONR'-, and -NR'-, where R' is alkyl, substituted alkyl, cycloalkyl, aryl, heteroaryl, heterocyclo; Ar is optionally substituted aryl, heteroaryl, or a protecting group;

 R_{70} and R_{70} are independently selected from the group consisting of hydrogen and methyl; R_{71} , R_{72} , R_{73} , and R_{74} are independently selected from the group consisting of hydrogen.

- 0 hydroxyl, and optionally substituted loweralkyl, cycloloweralkyl, cyclicaminoalkyl, alkylaminoalkyl, loweralkoxy, amino, alkylamino, alkylamino, alkylaminoyl, arylcarbonyl, aralkylcarbonyl, heteroarylcarbonyl, heteroaralkylcarbonyl, aryl and heteroaryl;
 - R_{75} and R_{78} are independently selected from the group consisting of hydrogen, halo, and optionally substituted loweralkyl, cycloalkyl, alkoxy, amino, aminoalkoxy, carbonyloxy, aminocarbonyloxy, alkylcarbonylamino, arylcarbonylamino, aralkylcarbonylamino, hetero-
- 5 aminocarbonyloxy, alkylcarbonylamino, arylcarbonylamino, aralkylcarbonylamino, heteroarylcarbonylamino, heteroaralkylcarbonylamino, cycloimido, heterocycloimido, amidino, cycloamidino, heterocycloamidino, guanidinyl, aryl, heteroaryl, heterocycloalkyl, heterocyclocarbonyloxy, heteroarylcarbonyloxy, and arylsulfonamido;

R₇₆ is selected from the group consisting of hydrogen, aryl, heteroaryl, substituted heteroaryl,

0 heterocyclyl, and substituted heterocyclyl;

R₇₇ is selected from the group consisting of hydrogen, hydroxy, halo, carboxyl, nitro, amino, amido, amidio, imido, cyano, sulfonyl, methanesulonyl, and substituted or unsubstituted alkyl, alkoxy, alkylcarbonyl, arylcarbonyl, aralkylcarbonyl, heteroarylcarbonyl, heteroaralkylcarbonyl, alkylcarbonyloxy, arylcarbonyloxy, aralkylcarbonyloxy, heteroarylcarbonyloxy, formyl, loweralkylcarbonyloxy, alkylaminocarbonyloxy, arylaminocarbonyloxy, formyl, loweralkylcarbonyl, loweralkoxycarbonyl, aminocarbonyl, aminoaryl, alkylsulfonyl, sulfonamido, aminoalkoxy, alkylamino, heteroarylamino, alkylcarbonylamino, alkylcarbonylamino, alkylcarbonylamino, heteroarylcarbonylamino, arylcarbonylamino, heteroarylcarbonylamino cycloamido, cycloamidio, heterocycloamidino, cycloalkyl, cycloimido, heterocycloimido, guanidinyl, aryl, heteroaryl, heterocyclo, heterocycloalkyl, arylsulfonyl and arylsulfonamido;

Anthraquinone compounds of the instant invention include, for example, compounds of Formula (XVI):

XVI

wherein.

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 R_{79} , R_{80} , R_{81} , and R_{82} may be the same or different and are independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonyl, aminocarbonyl, aminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups; and,

 R_{83} and R_{84} are taken together to form a substituted or unsubstituted 5-6 membered ring containing all carbon atoms or 1-2 heteroatoms selected from the group consisting of O, S, and N.

Quinoxaline compounds referred to throughout this application include tricyclic, partially unconjugated compounds optionally substituted with nitrogen heteroatoms as shown in the preferred quinoxaline embodiment (XVII) below:

wherein.

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J₁ is either C or N.

J₁' is selected from the group consisting of H, substituted aryl, unsubstituted aryl, substituted heteroaryl, and unsubstituted heteroaryl;

J₂ is either C or N,

J₂' is selected from the group consisting of H, substituted aryl, unsubstituted aryl, substituted heteroaryl, and unsubstituted heteroaryl;

 J_3 is selected from the group consisting of -CO-, -NH-, and -N=;

if J4 is -O- then J4' is absent; or,

if J_4 is =C- then J_4 ' is selected from the group consisting of H and substituted or unsubstituted alkyl, alkoxy, aryl, heteroaryl, heteroarylalkyl, arylalkyl, aminoalkyl, alkylamino, and alkylthio groups; and,

R₈₅, R₈₆, R₈₇, R₈₈, and R₈₉ may be the same or different and are independently selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonyl, aminocarbonyl, aminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylamino, heteroarylamino, heteroarylaminolkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups.

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Triazine compounds refer to substituted 6-membered heterocyclic groups with 3 nitrogen atoms distributed throughout the ring. The preferred embodiments of the instant invention include those shown in structures (XVIII), (XIX) and (XX) shown below:

wherein.

 R_{90} is selected from the group consisting of substituted or unsubstituted alkyl, alkenyl, akynyl, aryl, heteroarylalkyl, heteroarylalkyl, arylalkyl, and arylalkenyl; R_{91} and R_{93} are independently selected from the group consisting of H, and unsubstituted alkyl;

R₉₁ is aryl; preferably phenyl,

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wherein,

 R_{94} is selected from the group consisting of H, amino, alkyl, aminoalkyl, and halogen; R_{95} is selected from the group consisting of substituted or unsubstituted aryl, arylamino, arylalkylamino, heteroaryl, heteroarylamino, and heteroalkylamino;

 R_{96} and R_{97} are independently selected from the group consisting of H, halogen, and alkyl, preferably methyl; or,

 R_{96} may form a double bond with the nitrogen atom directly below it as indicated by the dashed line in the above structure; and.

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wherein.

 R_{98} is selected from the group consisting of H, substituted alkyl, and unsubstituted alkyl; preferably methyl.

R₉₉ is selected from the group consisting of H, substituted alkyl, and unsubstituted alkyl; preferably ethyl.

 R_{100} is selected from the group consisting of substituted or unsubstituted aryl, heteroaryl, alkoxyaryl, arylalkyl, and heteroarylalkyl.

Benzazole compounds as described throughout this application include compounds of the formula (XXI) as shown below:

wherein.

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A is selected from the group consisting of -O-, -S-, -NH-, and -NRs-:

W is selected from the group consisting of $-CH_{2^-}$, $-O_-$, $-S_-$, $-NH_-$, and $-NR_8$ -; R_7 is selected from the group consisting of carbocyclyl, unfused carbocyclylcarbocyclyl, substituted aryl, unsubstituted aryl, substituted heteroaryl, unsubstituted heteroaryl, substituted fused arylheteroaryl, unsubstituted fused arylheteroaryl, substituted unfused arylaryl;

 R_6 is selected from the group consisting of substituted or unsubstituted aryl, and heteroaryl; and,

R₈ is independently substituted or unsubstituted alkyl.

Pyrazalopyrimidine compounds as described throughout this application include compounds of formula (XXII) as shown below:

wherein.

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 R_{101} is selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, sulfonyl, aminosulfonyl, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups;

 R_{102} is selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups;

 R_{103} is selected from the group consisting of H, nitro, halogen, amino, hydroxy, cyano, carboxcyclic acid, trifluoromethyl, and substituted or unsubstituted alkyl, aryl, heteroaryl, alkoxy, alkylcarbonyl, alkylcarbonylamino, alkylaminocarbonyl, aminocarbonyl, arylalkoxy, heteroarylalkoxy, alkylamino, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, and carbocyclyl groups;

 R_{104} is selected from the group consisting of H and substituted or unsubstituted aryl, heteroaryl, arylalkoxy, heteroarylalkoxy, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclyl, heterocyclylalkoxy, heterocyclylalkyl, carbocyclylalkyl and carbocyclyl groups;

 R_{105} is selected from the group consisting of H and substituted or unsubstituted aryl, heteroaryl, arylalkoxy, heteroarylalkoxy, arylalkylamino, arylamino, heteroarylamino, heteroarylaminoalkyl, heterocyclylalkoxy, heterocyclylalkyl, carbocyclylalkyl and carbocyclyl groups;

wherein at least one of R_{104} and R_{105} is not H.

SMIP compounds identified by *in-vitro* (cellular or non-cellular assays) or *in-vivo* methods are thoroughly described in Methods 1 and 2 below.

Pharmaceutical compositions containing the compounds of the invention may be in any form suitable for the intended method of administration, including, for example, a solution, a suspension, or an emulsion. Liquid carriers are typically used in preparing solutions, suspensions, and emulsions. Liquid carriers contemplated for use in the practice of the present invention include, for example, water, saline, pharmaceutically acceptable organic solvent(s), pharmaceutically acceptable oils or fats, and the like, as well as mixtures of two or more thereof. The liquid carrier may contain other suitable pharmaceutically acceptable additives such as solubilizers, emulsifiers, nutrients, buffers, preservatives, suspending agents, thickening agents, viscosity regulators, stabilizers, and the like. Suitable organic solvents include, for example, monohydric alcohols, such as ethanol, and polyhydric alcohols, such as glycols. Suitable oils include, for example, soybean oil, coconut oil, olive oil, safflower oil, cottonseed oil, and the like. For parenteral administration, the carrier can also be an oily ester such as ethyl oleate, isopropyl myristate, and the like. Compositions of the present invention may also be in the form of microparticles, microcapsules, liposomal encapsulates, and the like, as well as combinations of any two or more thereof.

Other additives include immunostimulatory agents known in the art. Immunostimulatory oligonucleotides and polynucleotides are described in PCT WO 98/55495 and PCT WO 98/16247. U.S. Patent Application No. 2002/0164341 describes adjuvants including an unmethylated CpG dinucleotide (CpG ODN) and a non-nucleic acid adjuvant. U.S. Patent Application No. 2002/0197269 describes compositions comprising an antigen, an antigenic CpG-ODN and a polycationic polymer. Other immunostimulatory additives described in the art may be used, for example, as described in U.S. Patent No. 5,026,546; U.S. Patent No. 4,806,352; and U.S. Patent No. 5,026,543.

A controlled release delivery system may be used, such as a diffusion controlled matrix system or an erodible system, as described for example in: Lee, "Diffusion-Controlled Matrix Systems", pp. 155-198 and Ron and Langer, "Erodible Systems", pp. 199-224, in "Treatise on Controlled Drug Delivery", A. Kydonieus Ed., Marcel Dekker, Inc., New York 1992. The matrix may be, for example, a biodegradable material that can degrade spontaneously in situ and in vivo for, example, by hydrolysis or enzymatic cleavage, e.g., by proteases. The delivery system may be, for example, a naturally occurring or synthetic polymer or copolymer, for example in the form of a hydrogel. Exemplary polymers with cleavable linkages include polyesters, polyorthoesters, polyanhydrides, polysaccharides, poly(phosphoesters), polyamides, polyurethanes, poly(imidocarbonates) and poly(phosphazenes).

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The compounds of the invention may be administered enterally, orally, parenterally, sublingually, by inhalation spray, rectally, or topically in dosage unit formulations containing conventional nontoxic pharmaceutically acceptable carriers, adjuvants, and vehicles as desired. For example, suitable modes of administration include oral, subcutaneous, transdermal, transmucosal, iontophoretic, intravenous, intramuscular, intraperitoneal, intranasal, subdermal, rectal, and the like. Topical administration may also involve the use of transdermal administration such as transdermal patches or ionophoresis devices. The term parenteral includes subcutaneous injections, intravenous, intramuscular, intrasternal injection, or infusion techniques.

Injectable preparations, for example, sterile injectable aqueous or oleaginous suspensions may be formulated according to the known art using suitable dispersing or wetting agents and suspending agents. The sterile injectable preparation may also be a sterile injectable solution or suspension in a nontoxic parenterally acceptable diluent or solvent, for example, as a solution in 1,3-propanediol. Among the acceptable vehicles and solvents that may be employed are water, Ringer's solution, and isotonic sodium chloride solution. In addition, sterile, fixed oils are conventionally employed as a solvent or suspending medium. For this purpose any bland fixed oil may be employed including synthetic mono- or diglycerides. In addition, fatty acids such as oleic acid find use in the preparation of injectables.

Suppositories for rectal administration of the drug can be prepared by mixing the drug with a suitable nonirritating excipient such as cocoa butter and polyethylene glycols that are solid at ordinary temperatures but liquid at the rectal temperature and will therefore melt in the rectum and release the drug.

Solid dosage forms for oral administration may include capsules, tablets, pills, powders, and granules. In such solid dosage forms, the active compound may be admixed with at least one inert diluent such as sucrose lactose or starch. Such dosage forms may also comprise, as is normal practice, additional substances other than inert diluents, e.g., lubricating agents such as magnesium stearate. In the case of capsules, tablets, and pills, the dosage forms may also comprise buffering agents. Tablets and pills can additionally be prepared with enteric coatings.

Liquid dosage forms for oral administration may include pharmaceutically acceptable emulsions, solutions, suspensions, syrups, and elixirs containing inert diluents commonly used in the art, such as water. Such compositions may also comprise adjuvants, such as wetting agents, emulsifying and suspending agents, cyclodextrins, and sweetening, flavoring, and perfuming agents.

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As to the mode of administration, it should be emphasized that it is the combination of therapeutic agents that gives rise to its synergistic therapeutic effect no matter whether the first and the second agent are administered together or separately. Therefore, the two agents may be given together in a single dose or in separate ones with respect to space and time.

Effective amounts of the compounds of the invention generally include any amount sufficient to detectably treat viral infections.

Successful treatment of a subject in accordance with the invention may result in the inducement of a reduction or alleviation of symptoms in a subject afflicted with a medical or biological disorder to, for example, halt the further progression of the disorder, or the prevention of the disorder.

The amount of active ingredient that may be combined with the carrier materials to produce a single dosage form will vary depending upon the host treated and the particular mode of administration. It will be understood, however, that the specific dose level for any particular patient will depend upon a variety of factors including the activity of the specific compound employed, the age, body weight, general health, sex, diet, time of administration, route of administration, rate of excretion, drug combination, and the severity of the particular disease undergoing therapy. The therapeutically effective amount for a given situation can be readily determined by routine experimentation and is within the skill and judgment of the ordinary clinician.

The compounds of the present invention can also be administered in the form of liposomes. As is known in the art, liposomes are generally derived from phospholipids or other lipid substances. Liposomes are formed by mono- or multilamellar hydrated liquid crystals that are dispersed in an aqueous medium. Any non-toxic, physiologically acceptable and metabolizable lipid capable of forming liposomes can be used. The present compositions in liposome form can contain, in addition to a compound of the present invention, stabilizers, preservatives, excipients, and the like. The preferred lipids are the phospholipids and phosphatidyl cholines (lecithins), both natural and synthetic. Methods to form liposomes are known in the art. See, for example, Prescott, Ed., Methods in Cell Biology, Volume XIV, Academic Press, New York, N.W., p. 33 et seq (1976).

While the SMIP compounds of the invention can be administered as the sole active pharmaceutical agent, they can also be used in combination with one or more other agents used in the treatment of SARSs. Other representative agents useful in combination with the

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compounds of the invention for the treatment of viral infections include, for example, interferon, ribavirin, gancyclovir and the like.

When additional active agents are used in combination with the compounds of the present invention, the additional active agents may generally be employed in therapeutic amounts as indicated in the PHYSICIANS' DESK REFERENCE (PDR) 53rd Edition (1999), that is incorporated herein by reference, or such therapeutically useful amounts as would be known to one of ordinary skill in the art.

The compounds of the invention and the other therapeutically active agents can be administered at the recommended maximum clinical dosage or at lower doses. Dosage levels of the active compounds in the compositions of the invention may be varied so as to obtain a desired therapeutic response depending on the route of administration, severity of the disease and the response of the patient. The combination can be administered as separate compositions or as a single dosage form containing both agents. When administered as a combination, the therapeutic agents can be formulated as separate compositions that are given at the same time or different times, or the therapeutic agents can be given as a single composition.

Compounds of the present invention can be readily synthesized using the methods described herein, or other methods, that are well known in the art.

The compounds can be used in the form of salts derived from inorganic or organic acids. These salts include but are not limited to the following: acetate, adipate, alginate, citrate, aspartate, benzoate, benzenesulfonate, bisulfate, butyrate, camphorate, camphorsulfonate, digluconate, cyclopentanepropionate, dodecylsulfate, ethanesulfonate, glucoheptanoate; glycerophosphate, hemisulfate, heptanoate, hexanoate, fumarate, hydrochloride, hydrobromide, hydroiodide, 2-hydroxyethanesulfonate, lactate, maleate, methanesulfonate, nicotinate, 2-napthalenesulfonate, oxalate, pamoate, pectinate, persulfate, 3-phenylproionate, picrate, pivalate, propionate, succinate, tartrate, thiocyanate, p-toluenesulfonate and undecanoate. Also, the basic nitrogen-containing groups can be quaternized with such agents as loweralkyl halides, such as methyl, ethyl, propyl, and butyl chloride, bromides, and iodides; dialkyl sulfates like dimethyl, diethyl, dibutyl, and diamyl sulfates, long chain halides such as decyl, lauryl, myristyl and stearyl chlorides, bromides and iodides, aralkyl halides like benzyl and phenethyl bromides, and others. Water or oil-soluble or dispersible products are thereby obtained.

Examples of acids that may be employed to form pharmaceutically acceptable acid addition salts include such inorganic acids as hydrochloric acid, sulphuric acid and phosphoric acid and such organic acids as oxalic acid, maleic acid, succinic acid and citric acid. Basic

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addition salts can be prepared in situ during the final isolation and purification of the compounds of formula (I), or separately by reacting carboxylic acid moieties with a suitable base such as the hydroxide, carbonate or bicarbonate of a pharmaceutical acceptable metal cation or with ammonia, or an organic primary, secondary or tertiary amine. Pharmaceutical acceptable salts include, but are not limited to, cations based on the alkali and alkaline earth metals, such as sodium, lithium, potassium, calcium, magnesium, aluminum salts and the like, as well as nontoxic ammonium, quaternary ammonium, and amine cations, including, but not limited to ammonium, tetramethylammonium, tetraethylammonium, methylamine, dimethylamine, trimethylamine, triethylamine, ethylamine, and the like. Other representative organic amines useful for the formation of base addition salts include diethylamine, ethylenediamine, ethanolamine, diethanolamine, piperazine and the like.

Various compounds and methods of their synthesis are disclosed in international patent application Publication Nos. WO02/18327 (benzamide and pyridylamide based compounds); WO0222598, and WO02/18383 (ABIQ based compounds); and WO 02/81443 (pthalamide base compounds), that have been found within context of this invention to be useful for immune potentiation. The entire disclosure of these U.S. and international publications is incorporated herein by this reference. Other compounds or intermediates of interest in the present invention were purchased from commercially available sources using the following method: the chemical structure of interest was drawn into the ACD-SC database (from MDL Information Systems). A search of the following companies/institutions, among others, retrieved the identified compound's supplier and purchasing information: ASDI, ASINEX, BIONET, CHEMBRIDGE, CHEMDIV, CHEMEX, CHEMSTAR, COMGENEX, CSC, INTERBIOSCREEN, LABOTEST, MAYBRIDGE, MICROSOURCE/GENESIS, OLIVIA, ORION, PEAKDALE, RYAN SCIENTIFIC, SPECS, TIMTEC, U OF FLORIDA, and ZELINSKY.

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BENZAZOLE COMPOUNDS

Scheme 1

Compounds of the invention containing a benzimidazole core may be prepared using a number of methods familiar to one of skill in the art. In one method, suitably functionalized diamines may be coupled with various thioisocyanates to form the intermediate thioureas. Cyclization to form the benzimidazole moiety may be effected under known conditions such as with treatment carbodiimides or alkyl halides. Alternatively the diamines may be reacted

sequentially with carbonyl diimidazole and phosphoryl chloride followed by coupling with the appropriate amine.

Compounds containing the oxazole structure may similarly be prepared according to the methods above or according to other known general procedures. Haviv et. al. (J. Med. Chem. 1988, 31, 1719) describes a procedure for assembling oxazole cores wherein a hydroxy aniline is treated with ethyl potassium xanthate. The resulting sulfuryl benzoxazole may then be chlorinated and coupled with an amine.

Compounds containing a benzothiazole core may also be prepared according to known methods. An ortho-halothioisocyanate may be reacted with an amine to form a thiourea. Reduction with NaH then allows formation of the thiazole ring.

Benzothiazoles may generally be substituted in accordance with the present invention, such as through the following synthetic pathway:

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Synthesis of 4-[(2-{[4-chloro-3-(trifluoromethyl)phenyl]amino}-

1H-benzimidazol-6-yl)oxyl-N-methylpyridine-2-carboxamide
The compound 4-[(2-{[4-chloro-3-(trifluoromethyl)phenyl]amino}-1H-benzimidazol-6-yl)oxyl-N-methylpyridine-2-carboxamide (159322) was synthesized as follows:

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Step 1. Synthesis of 4-[(4-amino-3-nitrophenyl)oxy]-N-methylpyridine-2-carboxamide: A mixture containing 4-amino-3-nitrophenol (1eq) and potassium bis(trimethylsilyl)amide (2eq) was stirred in dimethylformamide for 2 hours at room temperature. To this mixture was added (4-chloro(2-pyridyl))-N-methylcarboxamide (1eq) and potassium carbonate (1.2eq) and stirred at 90°C for 3 days. The reaction mixture was then concentrated and partitioned between ethyl acetate and water. The organic layer was separated and washed with brine, dried, filtered, and concentrated in vacuum to give brown solid. Purification on silica gel (2% triethyl amine / 50% ethyl acetate in hexane) gave 4-[(4-amino-3-nitrophenyl)oxy]-N-methylpyridine-2-carboxamide as an orange solid. The product gave satisfactory NMR. HPLC, 3.39min; MS: MH⁺ = 289.

Step 2. Synthesis of 4-[(3,4-diaminophenyl)oxyl-N-methylpyridine-2-carboxamide: The mixture containing [4-(3-amino-4-nitrophenoxy)(2-pyridyl)]-N- in methanol with catalytic amount of 10%Pd/C was hydrogenated until disappearance of the yellow color to yield the product amine. HPLC, 2.5mins; MS: MH⁺ = 259.

Step 3. Synthesis of 4-[(2-{[4-chloro-3-(trifluoromethyl)phenyl]amino}-1H-benzimidazol-6-yl)oxyl-N-methylpyridine-2-carboxamide:

The mixture containing 4-[(3,4-diaminophenyl)oxy]-N-methylpyridine-2-carboxamide (1eq) and 4-chloro-3-(trifluoromethyl)benzeneisothiocyanate (1eq) in tetrahydrofuran was stirred at room temperature for 16 hours to give the corresponding thiourea. To the resulting mixture was added 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (2eq) and the mixture was stirred for another 10 hours. The mixture was concentrated and partitioned between ethyl acetate and water. The organic layer was washed with brine and dried. Purification on HPLC gave 4-[(2-

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{[4-chloro-3-(trifluoromethyl)phenyl]amino}-1H-benzimidazol-6-yl)oxy]-N-methylpyridine-2-carboxamide. MS: MH⁺ = 462

Synthesis of 4-({2-[(4-bromophenyl)amino]-1-methyl-

1H-benzimidazol-5-yl oxy)-N-methylpyridine-2-carboxamide

The compound 4-({2-[(4-bromophenyl)amino]-1-methyl-1H-benzimidazol-5-yl}oxy)-N-methylpyridine-2-carboxamide (161651) was synthesized as follows:

Step 1. Synthesis of 4-{[3-amino-4-(methylamino)phenyl]oxy}-N-methylpyridine-2-carboxamide: A solution of 4-[(4-amino-3-nitrophenyl)oxy]-N-methylpyridine-2-carboxamide (leq) in methylene chloride was treated with trifluoroacetic anhydride (leq) and stirred for 10 minutes at 0 °C. The mixture was quenched with satd. NaHCO₃ solution. The organic layer was separated and washed with water, brine, dried and evaporated. MS: MH*=385.2

To a solution of the trifluroacetamide (1eq) in a mixture of toluene, acetonitrile and sodium hydroxide solution (50%) was added benzyltrimethylammonium chloride (1eq) and dimethyl sulfate (1.2eq). The biphasic mixture was stirred overnight at room temperature and evaporated. The mixture was taken up in ethyl acetate, washed with water, brine, dried and evaporated. The crude product was purified by column chromatography eluting with 1:1 hexanes and ethylacetate followed by 2% triethylamine in 1:1 hexanes and ethyl acetate followed by 2% triethylamine in 1:1 hexanes and ethylacetate followed by 2% triethylamine in 1:1 hexanes and ethylacetate followed by 2% triethylamine in 1:1 hexanes and ethylacetate to afford N-methyl-4-{[4-(methylamino)-3-nitrophenyl]oxy}pyridine-2-carboxamide as a reddish orange solid. MS: MH $^+$ = 303.1.

The solution of nitromethylaniline in methanol was treated with 5% palladium on carbon and stirred under hydrogen atmosphere for 15 min. (until the disappearance of yellow coloration) at room temperature. The mixture was filtered and the filtrate was concentrated to provide 0.36 g of the diamine 4-{[3-amino-4-(methylamino)phenyl]oxy}-N-methylpyridine-2-carboxamide. MS: MH⁺ = 273.3.

Step 2. Synthesis of 4-{{2-[(4-bromophenyl)amino]-1-methyl-1H-benzimidazol-5-yl}oxy)-N-methylpyridine-2-carboxamide: A solution of the diamine 4-{[3-amino-4-(methylamino)phenyl]oxy}-N-methylpyridine-2-carboxamide (1eq) in methanol was treated with 4-bromophenylisothiocyanate (1eq) and stirred at 60 °C - 65°C for 2 hours. The reaction mixture was cooled down to room temperature and methyl iodide (1eq) was added and stirred overnight at 60°C. The reaction was cooled to room temperature, evaporated, taken up in ethyl -261-

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acetate, and washed with water and brine, dried, and evaporated under reduced pressure.

Column chromatography using a gradient solvent system of hexanes and ethyl acetate and either 1:1 methylene chloride and acetone or 5% methanol in methylene chloride yielded the product as a half white powder. MS: MH⁺=452.3

AMINOBENZIMIDAZOLYLQUINOLINONES

Compounds of structure I may be synthesized from simple starting molecules as shown in Schemes 1-4 and exemplified in the Examples. As shown in Scheme 1, compounds of structure I may generally be prepared using aromatic compounds substituted with amines and carboxylic acid groups.

Scheme 2.

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$$\begin{array}{c} \text{R} & \text{CO}_2\text{H} \\ \text{NH}_2 \end{array} + \begin{array}{c} \text{O} & \text{O} \\ \text{O} & \text{OMe} \end{array}$$

As shown in Scheme 2, a substituted aromatic compound such as a substituted or unsubstituted 2-aminobenzoic acid may be reacted with an acyl halide such as methyl 2-(chlorocarbonyl)acetate to produce an amide that will react with a substituted or unsubstituted 1,2-diaminobenzene. The resulting product is a 4-hydroxy-substituted compound of structure I. One skilled in the art will recognize that the procedure set forth in Scheme 1 may be modified to produce various compounds.

A method for preparing 4-amino substituted compounds of structure I is shown in Scheme 3. As shown in Scheme 3, aromatic compounds substituted with amine and nitrile groups may be used to synthesize 4-amino substituted compounds of structure I. A compound such as ethyl 2-cyanoacetate may be reacted with ethanol to produce ethyl 3-ethoxy-3-iminopropanoate hydrochloride. Subsequent reaction with a substituted or unsubstituted 1,2-phenylenediamine provides substituted or unsubstituted ethyl 2-benzimidazol-2-ylacetate. Reaction of a substituted or unsubstituted ethyl 2-benzimidazol-2-ylacetate with an aromatic

compound having an amine and nitrile group such as substituted or unsubstituted 2-aminobenzonitrile with a base such as lithium bis(trimethylsilyl)amide or a Lewis acid such as tin tetrachloride provides the substituted or unsubstituted 4-amino substituted compound of structure I.

Scheme 4 illustrates a general synthetic route that allows for the synthesis of 4-dialkylamino and 4-alkylamino compounds of structure I. An inspection of Scheme 3 shows that 4-hydroxy substituted compounds of structure I may be converted into the 4-chloro derivative by reaction with phosphorous oxychloride or thionyl chloride. The 4-chloro derivative may then be reacted with an alkylamine or dialkylamine to produce the corresponding 4-alkylamino or 4-dialkylamino derivative. Deprotection affords the final 4-alkylamino or 4-dialkylamino compounds of structure I. Other groups that may be reacted with the 4-chloro derivative in this manner include, but are not limited to, ROH, RSH, and CuCN.

SnCl

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Scheme 4

As shown in Scheme 5, the synthesis of compounds of structure I having a H, alkyl group, aryl group, or heterocyclyl group in the 4-position may be accomplished using a substituted or unsubstituted 2-benzimidazol-2-ylacetate prepared as shown in Schemes 3 and 4.

Scheme 5.

R" = H, alkyl aryl, heterocyclyl

THIOSEMCARBAZONES

A solution of aldehyde (1.0 equiv.) and thiosemicarbazide (1.05 equiv.) in acetic acid was stirred overnight. Excess of acetic acid was removed to give a residue, that was washed with ethanol, or purified by preparative-HPLC to give the thiosemicarbazone.

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Scheme 7

A solution of aldehyde (1.0 equiv.), thiosemicarbazide (1.05 equiv.) and acetic acid (0.1 equiv.) in methanol was stirred overnight. Methanol was removed to give a residue, that was worked up as in Scheme 6.

5 Scheme 8

To a solution of {[[(1E)-1-aza-2-(4-fluoro-3-nitrophenyl)vinyl]amino}-aminomethane-1-thione in ethanol was added an arylamine (2.1 equiv.). The solution was stirred at room temperature until the starting fluoride disappeared. The solution was purified to the product.

Scheme 9

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A mixture of 4-(diethylamino)-2-hydroxybenzaldehyde (1 equiv.), benzylic bromide (1.2 equiv.) and powder potassium carbonate in ethanol was stirred at room temperature for 2 days. Ethanol was removed, and the residue was dissolved in ethyl acetate and water. The organic layer was washed with aqueous NaHCO₃ and brine, dried over Na₂SO₄., and concentrated. The residue was purified on silica gel eluting with ethyl acetate/hexane to give 4-(diethylamino)-2-benzoxylic-benzaldehyde.

The aldehydes were converted to thiosemicarbazones according to Scheme 7.

Scheme 10

A solution of 3,4-difluorobenzenecarbonitrile (1 equiv.), amine (1.5 equiv.) and DIEA (2 equiv.) in NMP was heated in a Smith Microwave (Personal Chemistry) for 30 minutes. The reaction mixture was purified on silica gel to give 4-substituted 3-fluorobenzenecarbonitrile.

To a solution of nitrile in toluene at -78 °C was added DIBAL-H (1 M in toluene, 1.5 equiv.). The reaction mixture was warmed to rt, and stirred for 16 h, and quenched with methanol/ethyl acetate/brine (1:1:4). After being stirred at rt for 30 min, the solution was extracted with ethyl acetate (3x). The combined organic layers were washed with aqueous NaHCO₃, brine and concentrated. The aldehyde was purified on silica gel or directly converted to thiosemicarbazones (Scheme 7).

10 Scheme 11

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A solution of 2,4,5-trifluorobenzenecarbonitrile (1 equiv.) and 4-arylpiperazine (1.2 equiv.) and DIEA (1.2 equiv.) in THF was heated at 80 °C for 2 hours. The mixture was purified on silica gel to give 4-substituted 2,5-difluorobenzenecarbonitrile.

Scheme 12

To an alcohol (1.0 equiv) was added potassium t-butoxide in THF (1 M, 1.1 equiv). After 5 minutes, the solution was added to a solution of 4-N-substituted-2,5-diffluorobenzenecarbonitrile (1 equiv.) in THF. The reaction mixture was stirred at rt overnight and quenched with aqueous ammonium chloride. The aqueous layer was extracted with ethyl acetate (3x). The combined organic layers were washed with brine, and concentrated to give a residue, that was purified to give 4-N-substituted-2-O-substituted-5-fluorobenzenecarbonitrile.

4-N-substituted-2-O-substituted-5-fluorobenzenecarbonitrile was reduced with DIBAL-H to give a 4-N-substituted-2-O-substituted-5-fluorobenzaldehyde according to procedure in Scheme 10.

The aldehyde was converted to the corresponding thiosemicarbazone using Scheme 7. Scheme 13

A solution of 4-N-substituted-2,5-difluorobenzenecarbonitrile (1 equiv.), amine (1.5 equiv.) and DIEA (2 equiv.) in NMP was heated in a Smith Microwave (Personal Chemistry) for 30 minutes. The reaction mixture was purified on silica gel to give 4-N-substituted-2-N-substituted-5-fluorobenzenecarbonitrile.

4-N-substituted-2-N-substituted-5-fluorobenzenecarbonitrile was reduced with DIBAL-H according to procedure described in Scheme 10 to give 4-N-substituted-2-N-substituted-5-fluorobenzaldehyde.

 $\label{preparation} Preparation of a mino \{3-[5-(3-chlorophenyl)(2-furyl)](2-pyrazolinyl)\} methane-1-thione$

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To a solution of 5-(3-chlorophenyl)furan-2-carbaldehyde (1.0 equiv.) in THF at 0 $^{\circ}$ C was added MeMgBr in ether (3.0 equiv.) and stirred for 45 min. The reaction was quenched with water, diluted with ether and filtered through Celite. The organic layer was separated and washed with brine, dried over MgSO₄, and concentrated to give the 1-[5-(3-chlorophenyl)-2-furyl]ethan-1-ol.

To a solution of secondary alcohol(1.0 equiv.) in CH_2Cl_2 was added MnO_2 (10 equiv.). The reaction was stirred overnight, filtered through Celite, and concentrated to give 1-[5-(3-chlorophenyl)-2-furyl]ethan-1-one.

To a mixture of ketone (1.0 equiv.), paraformaldehyde (2.0 equiv.), and dimethylamine hydrochloride (2.0 equiv) and molecular sieves in ethanol was added concentrated hydrochloric acid (cat.). The reaction was refluxed overnight under nitrogen and the concentrated. A few drops of HCl was added, and the mixture was worked up with DCM and water. The organic layer was discarded. The aqueous layer was adjusted to basic and extracted with DCM (3x). The organic layer was washed with brine, dried over MgSO₄, and concentrated to yield 3-(dimethylamino)-1-[5-(3-chlorophenyl)(2-furyl)]propan-1-one.

Thiosemicarbazide (1.0 equiv.) was dissolved in MeOH upon heating under nitrogen. Aqueous sodium hydroxide (6 M, 9.0 equiv.) was added to the reaction. A methanol solution of 3-(dimethylamino)-1-[5-(3-chlorophenyl)(2-furyl)]propan-1-one (1.0 equiv) was then added dropwise to the reaction mixture. The solvent was removed and the residue was dissolved in DCM and washed with water, brine, dried over MgSO₄, and concentrated. The final compound was purified by preparative-HPLC to give amino {3-[5-(3-chlorophenyl)(2-furyl)](2-purgazolinyl)]methano 1 thicary LCOM are 10.000 of 10.000 preparative properties of 10.0000 preparative properties of 10.000 preparative properties of 10.000 prepara

25 pyrazolinyl)}methane-1-thione; LC/MS m/z 306.2 (MH+); Rt =3.06 minutes . Scheme 14

To a solution of 4-pyridylmethylamine (1.0 equiv.) and triethylamine (2.0 equiv.) in $CHCl_3$ was added CS_2 (1.0 equiv.)) and stirred overnight. The reaction was cooled to 0 °C and ethyl chloroformate (1.0 equiv.) was added dropwise. The reaction was stirred for 15 min at 0 °C and then stirred at room temperature for 2 hrs followed by addition of (tert-

butyl)oxycarbohydrazide (1.2 equiv.). After stirring for an addition hour the mixture was washed with aqueous citric acid (5%), saturated NaHCO₃, brine, dried over MgSO₄, and concentrated. The desired Boc protected thiosemicarbazide was purified using column - chromatography.

To a solution of Boc protected thiosemicarbazide (1.0 equiv.) dissolved in DCM was added HCl in dioxane (2M, 8.3 equiv.) and stirred for 15 min. MeOH is then added to dissolve the precipitate, followed by addition of the furfural, and small amount of acetic acid (0.5 mL). The mixture is stirred overnight and the solvents are removed to give a residue purified by preparative-HPLC to give the thiosemicarbazone.

Synthesis of 4-[4-(4-methylpiperazin-1-yl)phenoxymethyl]benzaldehyde

To a solution of 4-piperazin-1-yl phenol (1 equivalent) in CHCl₃, cooled to 0 °C, was added di-t-butyl dicarbonate (1 equivalent) in CHCl₃ drop-wise. The solution was stirred at 0 °C for 1 hour before removing from the cold bath and stirring at ambient temperatures for 18 hours. The organic solution was washed aqueous NaHCO₃ and brine dried over MgSO₄ and concentrated the crude material was used without purification.

A solution of the resulting 4-(1-BOC-piperazin-4-yl)phenol (1 equivalent) in dry CH₃CN was slowly added drop-wise to a slurry of NaH (1 equivalent) in dry CH₃CN at room temperature under N₂. The slurry was stirred at room temperature for 2 hours before the solids were filtered and washed with Et₂O.

Sodium 4-(1-BOC-piperazin-4-yl)phenoxide (1 equivalent) and methyl 4-bromomethylbenzoate (1 equivalent) were combined in dry acetone and heated to reflux at 60 °C for 18 hours. The slurry was filtered and the filtrate was then concentrated to provide the crude methyl 4-[4-(1-BOC-piperazin-4-yl)phenoxymethyl]benzoate, that was used without purification.

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having sequence identity to an amino acid sequence selected from the group consisting of the sequences shown in Figure 127, and in particular SEQ ID NO^S: 10506 to 10570.

The invention also provides fragments of amino acid sequences encoded by SEQ ID NO: 10505. The invention also provides fragments of amino acid sequences selected from the group consisting of SEQ ID NO $^{\rm S}$: 10506 to 10570. In one embodiment, the fragment does not consist entirely of a known amino acid sequence of a SARS virus or a known amino acid sequence of a coronavirus.

In one embodiment, the invention includes a polypeptide comprising an amino acid sequence from the 5'3' Frame 3 of Figure 127. Some encoded open reading frames within this translation are: SEQ ID NO: 10533; SEQ ID NO: 10571; SEQ ID NO: 10572; SEQ ID NO: 10573; SEQ ID NO: 10574.

The invention includes a polypeptide sequence comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10533, SEQ ID NO: 10571, SEQ ID NO: 10572, SEQ ID NO: 10573 and SEQ ID NO: 10574. The invention includes a polypeptide having sequence identity to the amino acid sequence selected from the group consisting of SEQ ID NO: 10533, SEQ ID NO: 10571, SEQ ID NO: 10572, SEQ ID NO: 10573 and SEQ ID NO: 10574. The invention includes a fragment of a polypeptide sequence comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10533, SEQ ID NO: 10571, SEQ ID NO: 10572, SEQ ID NO: 10573, SEQ ID NO: 10573, SEQ ID NO: 10573, SEQ ID NO: 10574.

Partial BLAST results of SEQ ID NO: 10533 against GenBank are given below:

>gi|7739601|gb|AAF68926.1|AF207902_11 nucleocapsid protein [murine hepatitis virus strain ML-11] Length = 451

Score = 147 bits (370), Expect = 3e-34 Identities = 102/252 (40%), Positives = 137/252 (54%), Gaps = 18/252 (7%)

Query: 49 SWFTALTQHGK-EELRFPRGGGVPINTNSGPDDQIGYYRRATRR-VRGGDGKMKELSPRW 106
SWF+ +TQ K +E +F +GQGVPI + +Q GY+ R RR + DG+ K+L PRW
Sbjct: 63 SWFSGITQFQKGKEFQFAQGGGVPIASGIPASEQKGYWYRHNRRSFKTPDGQHKQLLPRW 122

Query: 107 YFYYLGTGPEASLFYGANKEGIVWVATEGALNTPKDHIGTRNPNNNAATVLQLFQGTTLF 166
YFYYLGTGP A YG + EG+VWVA++ A PAPLLLA A YG + BAPLLLA A YG + BAPLLA A YG + BAPLA A YG + BAPLLA A YG + BAPLA A YG + BAPLLA A YG + BAPLA A

Query: 167 KGFYAEGSRGGSQASSRSSSRSRGNSRNSTPGSSRGNSPARMASGGGETALALLLDRLN 226 FGFY EGS + AS S N SS PA +A L+L +L

Sbjet: 183 QGFYVEGSGRSAPASRSGSRSQSRGPNNRARSSSNQRQPASAVKPDMAEEIAALVLAKLG 242
Query: 227 QLESKVSGKGQQQQGQTVTKKSAAEASK----KPRQKRTATKQYNVTQAFGRRGPEQTQG 282

Sbjct: 243 K----DAGQPKQ---VTKQSAKEVRQKILTKPRQKRTFKQCPVQQCPGKRGPNQ--- 290

Query: 283 NFGDQDLIRQGT 294 NFG ++++ GT

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_2004092360A2_l_>

NFG ++++ GT Sbjct: 291 NFGGSEMLKLGT 302

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>gi|3132999|gb|AAC16422.1| nucleocapsid protein [murine hepatitis virus
       strain 21
 5
                 Length = 451
        Score = 147 bits (370), Expect = 3e-34
        Identities = 102/252 (40%), Positives = 137/252 (54%), Gaps = 18/252 (7%)
10
                 SWFTALTOHGK-EELRFPRGOGVPINTNSGPDDOIGYYRRATRR-VRGGDGKMKELSPRW 106
       Ouerv: 49
                  SWF+ +TQ K +E +F +GQGVPI + +Q GY+ R RR + DG+ K+L PRW
                 SWFSGITQFQKGKEFQFAQGQGVPIASGIPASEQKGYWYRHNRRSFKTPDGQHKQLLPRW 122
       Sbict: 63
       Query: 107 YFYYLGTGPEASLPYGANKEGIVWVATEGALNTPKDHIGTRNPNNNAATVLOLPOGTTLP 166
15
                 YFYYLGTGP A YG + EG+VWVA++ A
                                                    + R+P+++ A + GT LP
       Sbict: 123 YFYYLGTGPHAGAEYGDDIEGVVWVASQQADTKTTADVVERDPSSHEAIPTKFAPGTVLP 182
       Query: 167 KGFYAEGSRGGSOASSRSSSRSRGNSRNSTPGSSRGNSPARMASGGGETALALLLLDRLN 226
                 +GFY EGS + AS
                                 S
                                            N
                                                SS
                                                      PA
                                                                 +A L+L +L
20
       Sbjct: 183 QGFYVEGSGKSAPASRSGSRSQSRGPNNRARSSSNORQPASAVKPDMAEEIAALVLAKLG 242
       Query: 227 QLESKVSGKGQQQQGTVTKKSAAEASK----KPROKRTATKQYNVTQAFGRRGPEQTQG 282
                          GQ +Q VTK+SA E + KPRQKRT KQ V O FG+RGP O
       Sbjct: 243 K-----DAGQPKQ---VTKQSAKEVRQKILTKPRQKRTPNKQCPVQQCFGKRGPNQ--- 290
25
       Query: 283 NFGDQDLIRQGT 294
                 NFG ++++ GT
       Sbjct: 291 NFGGSEMLKLGT 302
30
       gi | 127877 | sp | P03417 | NCAP_CVMJH
                                      Nucleocapsid protein
       gi 74859 pir VHIHMJ
                                nucleocapsid protein - murine hepatitis virus
       (strain JHM)
35
        gi|58973|emb|CAA25497.1| nucleocapsid protein [Murine hepatitis virus]
                Length = 455
        Score = 146 bits (369), Expect = 4e-34
       Identities = 110/254 (43%), Positives = 142/254 (55%), Gaps = 22/254 (8%)
10
       Query: 49 SWFTALTQHGK-EELRFPRGQGVPINTNSGPDDOIGYYRRATRR-VRGGDGKMKELSPRW 106
                 SWF+ +TO K +E +F +GOGVPI
                                                Q GY+ R RR .+ DG+ K+L PRW
      Sbjct: 67 SWFSGITQFQKGKEFQFAQGQGVPIANGIPASQQKGYWYRHNRRSFKTPDGQQKQLLPRW 126
15
       Query: 107 YFYYLGTGPEASLPYGANKEGIVWVATEGALNTPKDHIGTRNPNNNAATVLQLPQGTTLP 166
                 YFYYLGTGP A YG + EG+VWVA++ A
                                                    I R+P+++ A + GT LP
       Sbjct: 127 YFYYLGTGPYAGAEYGDDIEGVVWVASQQAETRTSADIVERDPSSHEAIPTRFAPGTVLP 186
       Ouery: 167 KGFYAEGSRGGSQASSRSSSR--SRGNSRNSTPGSSRGNSPARMASGGGETALALLLLDR 224
50
                 +GFY EGS G S +SRS SR SRG N
                                                 SS
                                                       PA
       Sbjct: 187 QGFYVEGS-GRSAPASRSGSRPOSRG-PNNRARSSSNOROPASTVKPDMAEETAALVLAK 244
      Ouery: 225 LNQLESKVSGKGQQQQGQTVTKKSAAEASK----KPRQKRTATKQYNVTQAFGRRGPEQT 280
                 T. +
                            GQ +Q VTK+SA E + KPROKRT KO V O FG+RGP O
55
      Sbjct: 245 LGK-----DAGQPKQ---VTKQSAKEVRQKILNKPRQKRTPNKQCPVOQCFGKRGPNO- 294
      Query: 281 QGNFGDQDLIRQGT 294
                   NFG ++++ GT
      Sbict: 295 --NFGGPEMLKLGT 306
'n
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5		hepati	tis 6934 tis	6 gb AAF19389.1 AF201929_7 virus strain 2 8 gb AAF69338.1 AF208066_11 virus Length = 451	nucleocapsid	-	[murine
10		Score = 146 bits (368), Expect = 5e-34 Identities = 102/252 (40%), Positives = 137/252 (54%), Gaps = 18/252 (7%)					
	•	Query: Sbjct:		SWFTALTQHGK-EELRFPRGQGVPINTNSG SWF+ +TQ K +E +F +GQGVPI + SWFSGITQFQKGKEFQFAQGQGVPIASGIP	+Q GY+ R RR +	DG+ K+L	PRW -
15				YFYYLGTGPEASLPYGANKEGIVWVATEGA YFYYLGTGP A YG + EG+VWVA++ A YFYYLGTGPHAGAEYGDDIEGVVWVASQQA	+ R+P+++	A + GT	T.D
20		* *		KGFYAEGSRGGSQASSRSSRSRGNSRNST +GFY EGS + AS S N QGFYVEGSGRSAPASRSGSRSQSRGPNNRA	SS PA	+A L+L -	+I.
25		Query:	227	QLESKVSGKGQQQGGTVTKKSAAEASK + GQ +Q VTK+SA E + KDAGQPKQVTKQSAKEVRQKI	KPRQKRTATKQYNVT KPROKRT KO V	QAFGRRGPEQ	PQG 282
		Query:	283	NFGDQDLIRQGT 294 NFG ++++ GT	DIAPROKITPHKOCPVC	QCFGKRGPNQ-	290
30		Sbjct:	291	NFGGSEMLKLGT 302			
35		<pre>>gi 21734854 gb AAM77005.1 AF481863_7 phosphorylated nucleocapsid protein N [porcine hemagglutinating encephalomyelitis virus] Length = 449</pre>					
	Score = 145 bits (366), Expect = 8e-34 Identities = 107/253 (42%), Positives = 145/253 (57%), Gaps = 18/253						
40	٠,	Query: Sbjct:		SWFTALTQHGK-EELRFPRGQGVPINTNSG SWF+ +TQ K +E F GQGVPI SWFSGITQFQKGKEFEFAEGQGVPIAPGVP	+ GY+ R RR +	DG ++L 1	PRW
45		Query:	107	YFYYLGTGPEASLPYGANKEGIVWVATEGA YFYYLGTGP A YG + +G+ WVA+ A YFYYLGTGPHAKHQYGTDIDGVFWVASNQA	- -LNTPKDHIGTRNPNNN - +NTP D I R+P+++	AATVLQLPQG	TTL 165
50 55				PKGFYAEGSRGGSQASSRSSSRSRGNSRNS	TPGSSRGNSPARMASGG	GETALALLLLI	ORL 225
				PQGYYIEGS-GRSAPNSRSTSRA-PNRAPS.	AGSRSRANSGNRTSTPG	VTPDMAI	
				$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	KPROKR+ KO V	O FG+RGP ()
				GNFGDQDLIRQGT 294 NFG ++++ GT . -NFGGGEMLKLGT 305			
60		_2,00.					

```
>gi | 23295765 | gb | AAL80036.1 |
                                              nucleocapsid
                                                               protein
                                                                          [porcine
      hemagglutinating encephalomyelitis virus]
                Length = 449
5
        Score = 145 bits (365). Expect = 1e-33
       Identities = 107/253 (42%), Positives = 145/253 (57%), Gaps = 18/253 (7%)
                 SWFTALTOHGK-EELRFPRGOGVPINTNSGPDDOIGYYRRATRR-VRGGDGKMKELSPRW 106
                 SWF+ +TO K +E F GOGVPI
                                                 + GY+ R RR + DG ++L PRW
10
       Sbjct: 64 SWFSGITOFOKGKEFEFAEGOGVPIAPGVPSTEAKGYWYRHNRRSFKTADGNOROLLPRW 123
       Query: 107 yfyylgtgpeaslpygankegivwvatega-lntpkdhigtrnpnnnaatvlolpogttl 165
                 YFYYLGTGP A
                              YG + +G+ WVA+ A +NTP D I R+P+++ A
       Sbict: 124 YFYYLGTGPHAKDOYGTDIDGVFWVASNOADINTPAD-IVDRDPSSDEAIPTRFPPGTVL 182
15
       Ouery: 166 PKGFYAEGSRGSQASSRSSSRSRGNSRNSTPGSSRGNSPARMASGGGETALALLLLDRL 225
                  P+G+Y EGS G S +SRS+SR+ N S · SR NS · R ++ G
                                                                    +A
                                                                          D++
       Sbjct: 183 POGYYIEGS-GRSAPNSRSTSRA-PNRAPSAGSRSRANSGNRTSTPGVTPDMA----DOI 236 .
20
       Ouery: 226 NOLESKYSGKGOOOGGTVTKKSAAEASK----KPROKRTATKOYNVTOAFGRRGPEOTO 281
                         GK
                               + Q VTK++A E +
                                                 KPROKR+ KO V O FG+RGP O
                   т.
       Sbjct: 237 ASLVLAKLGK-DATKPQQVTKQTAKEVRQKILNKPRQKRSPNKQCTVQQCFGKRGPNQ-- 293
       Ouery: 282 GNFGDQDLIROGT 294
25
                  NFG ++++ GT
       Sbict: 294 -NFGGGEMLKLGT 305
```

These results indicate that SEQ ID NO: 10533 has functional similarities to a coronavirus nucleocapsid protein.

In one embodiment, the invention comprises an amino acid sequence from the 5'3' Frame 1 of Figure 127 e.g. SEQ ID NO^S: 10506-10514. Some encoded open reading frames within this region are SEQ ID NO^S: 10575 to 10578.

Accordingly, the invention includes a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 10575, SEQ ID NO: 10576, SEQ ID NO: 10577 and SEQ ID NO: 10578. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to a sequence selected from the group consisting of SEQ ID NO: 10097, SEQ ID NO: 10576, SEQ ID NO: 10577 and SEQ ID NO: 10578. The invention includes a fragment of a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 10097, SEQ ID NO: 10576, SEQ ID NO: 10577 and SEQ ID NO: 10578.

In one embodiment, the invention includes a polypeptide comprising an amino acid sequence from the 3'5' Frame 2 of Figure 127 e.g. SEQ ID NO⁵: 10547-10559. An open reading frame within this region is SEO ID NO: 10579.

The invention includes a polypeptide comprising an amino acid sequence of SEQ ID NO: 10579. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 10579. The invention includes a fragment of a polypeptide comprising an amino acid sequence of SEQ ID NO: 10579.

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The invention also includes polynucleotide sequences which can be used as probes for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified in Table 33. The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified in Table 33.

The invention includes a polynucleotide sequence comprising SEQ ID NO: 11323. A polypeptide encoded by SEQ ID NO: 11323 is SEQ ID NO: 11324.

The invention includes a polypeptide comprising SEQ ID NO: 11324, sequence having sequence identity to SEQ ID NO: 11324 and fragments of SEQ ID NO: 11324. The invention includes a fragment of SEQ ID NO: 11324, wherein said polypeptide fragment begins with a Methionine.

Accordingly, the invention includes a polynucleotide sequence comprising SEQ ID NO: 11323. It also provides polynucleotide sequences having sequence identity to SEQ ID NO: 11323. The invention also provides for polynucleotide sequences comprising fragments of SEQ ID NO: 11323. In one embodiment, the polynucleotide fragment does not consist entirely of a known SARS polynucleotide sequence or a known coronavirus polynucleotide sequence.

The invention includes an amino acid sequence encoded by the polynucleotide sequence SEQ ID NO: 11323, including the amino acid sequence of SEQ ID NO: 11324.

The invention also provides amino acid sequences having sequence identity to an amino acid sequence encoded by SEQ ID NO: 11323. The invention provides amino acid sequences having sequence identity to SEQ ID NO: 11324.

The invention provides fragments of amino acid sequences encoded by SEQ ID NO: 11323. The invention also provides fragments of amino acid sequences of SEQ ID NO: 11324. In one embodiment, the fragment does not consist entirely of a known SARS amino acid sequence or a known coronavirus amino acid sequence.

The invention also includes polynucleotide sequences which can be used as probes for diagnostic reagents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample. The invention includes a polynucleotide sequence comprising one or more of the primer sequences identified as SEQ ID NO^S: 11325-11440 (left part) and SEQ ID NO^S: 11441-11551 (right part). The invention further includes polynucleotide sequence comprising the complement of one or more of the primer sequences identified as SEQ ID NO^S: 11325-11551.

The invention includes a polypeptide comprising SEQ ID NO: 11552. The SARS virus contains polymorphism at the Isoleucine residue Ile-324. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11552, wherein

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said polypeptide includes an amino acid sequence selected from the group consisting of YSYAI (SEQ ID NO: 11553), SYAIH (SEQ ID NO: 11554), YAIHH (SEQ ID NO: 11555), IHHDK (SEQ ID NO: 11556), SYAI (SEQ ID NO: 11557), YAIH (SEQ ID NO: 11558), AIHH (SEQ ID NO: 11559), IHHD (SEQ ID NO: 11560), YAI, AIH, and IHH. The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11552, wherein said fragment includes an amino acid sequence selected from the group consisting of YSYAI (SEQ ID NO: 11553), SYAIH (SEQ ID NO: 11554), YAIHH (SEQ ID NO: 11555), IHHDK (SEQ ID NO: 11556), SYAI (SEQ ID NO: 11557), YAIH (SEQ ID NO: 11558), AIHH (SEQ ID NO: 11559), IHHD (SEQ ID NO: 11560), YAI, AIH, and IHH.

The invention includes a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 11561 and SEQ ID NO: 11562. The invention includes a fragment of a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 11561 and SEQ ID NO: 11562.

The invention includes a diagnostic kit comprising a polypeptide comprising at least one of the amino acid sequences selected from the group consisting of SEQ ID NO⁸: 11561 and 11562. The invention includes a diagnostic kit comprising a polypucleotide sequence encoding a polypeptide comprising at least one of the amino acid sequences selected from the group consisting of SEQ ID NO⁸: 11561 and 11562. The invention includes an immunogenic composition comprising a polypeptide comprising at least one of the amino acid sequences selected from the group consisting of SEQ ID NO⁸: 11561 and 11562. The invention includes an antibody which recognizes a polypeptide comprising at least one of the amino acid sequences selected from the group consisting of SEO ID NO⁸: 11561 and 11562.

The invention includes a polynucleotide sequence SEQ ID NO: 11563 or a fragment thereof or a sequence having sequence identity thereto. Polypeptide sequences which can be translated from SEQ ID NO: 11563 are shown in Figure 128. The constituent amino acid sequences from Figure 128, having at least 4 amino acids, are listed as SEQ ID NO⁸: 11564 to 11617.

The invention includes a polypeptide sequence selected from the group consisting of the sequences of Figure 128, or a fragment thereof or a sequence having sequence identity thereto e.g. SEQ ID NOS: 11563 to 11617.

A polypeptide sequence within SEQ ID NO: 11600 is SEQ ID NO: 11618. The invention includes a polypeptide comprising SEQ ID NO: 11618, or a fragment thereof or a sequence having sequence identity thereto.

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A polypeptide sequence within SEQ ID NO: 11602 is SEQ ID NO: 11641. The invention includes a polypeptide comprising SEQ ID NO: 11641, or a fragment thereof or a sequence having sequence identity thereto.

A polypeptide sequence within SEQ ID NO: 11609 is SEO ID NO: 11619.

The invention includes a polynucleotide encoding (i) an amino acid sequence selected from the group consisting of: (1) the amino acid sequences of Figure 128, and in particular SEQ ID NOS: 11564-11617; (2) SEQ ID NO: 11618; and (3) SEQ ID NO: 11619, or (ii) a fragment thereof. The invention includes a diagnostic kit comprising a one or more of these proteins. The invention includes a diagnostic kit comprising a polynucleotide sequence encoding one or more of these polypeptide sequences. The invention includes an antibody which recognizes one or more of the polypeptide sequences.

The SARS virus may contain polymorphism at isoleucine residue IIe-326 in SEQ ID NO: 11620 (Chi-PEP3). The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11620, wherein said polypeptide includes an amino acid sequence selected from the group consisting of YAIHH (SEQ ID NO: 11621) and YAIHH (SEQ ID NO: 11622). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11620, wherein said fragment includes an amino acid sequence selected from the group consisting of YAIHH (SEQ ID NO: 11621) and YAIHH (SEQ ID NO: 11622).

The SARS virus may contain polymorphism at glutamine residue Gln-830 in SEQ ID NO: 11620. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11620, wherein said polypeptide includes an amino acid sequence selected from the group consisting of ASQAW (SEQ ID NO: 11623) and ASRAW (SEQ ID NO: 11624). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11620, wherein said fragment includes an amino acid sequence selected from the group consisting of ASQAW (SEQ ID NO: 11623) and ASRAW (SEQ ID NO: 11624).

The SARS virus may contain polymorphism at aspartic acid residue Asp-935 in SEQ ID NO: 11620. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11620, wherein said polypeptide includes an amino acid sequence selected from the group consisting of DADST (SEQ ID NO: 11625) and DAYST (SEQ ID NO: 11626). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11620, wherein said fragment includes an amino acid sequence selected from the group consisting of DADST (SEQ ID NO: 11625) and DAYST (SEQ ID NO: 11626).

The SARS virus may contain polymorphism at serine residue Ser-577 in SEQ ID NO: 11627 (Chi-PEP4). The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11627, wherein said polypeptide includes an amino

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acid sequence selected from the group consisting of PCSFG (SEQ ID NO: 11628) and PCAFG (SEQ ID NO: 11629). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11627, wherein said fragment includes an amino acid sequence selected from the group consisting of PCSFG (SEQ ID NO: 11628) and PCAFG (SEQ ID NO: 11629).

The SARS virus may contain polymorphism at valine residue Val-68 in SEQ ID NO: 11630 (Chi-PEP8). The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11630, wherein said polypeptide includes an amino acid sequence selected from the group consisting of LAVVY (SEQ ID NO: 11631) and LAAVY (SEQ ID NO: 11632). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11630, wherein said fragment includes an amino acid sequence selected from the group consisting of LAVVY (SEQ ID NO: 11631) and LAAVY (SEQ ID NO: 11632).

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The SARS virus may contain polymorphism at isoleucine residue Ile-50 in SEQ ID NO: 11633 (Chi-PEP13). The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11633, wherein said polypeptide includes an amino acid sequence selected from the group consisting of NNIAS (SEQ ID NO: 11634) and NNIAS (SEQ ID NO: 11635). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11633, wherein said fragment includes an amino acid sequence selected from the group consisting of NNIAS (SEQ ID NO: 11635).

The SARS virus may contain a polymorphism at Serine residue Ser-943 in SEQ ID NO: 11636. The invention includes a polypeptide comprising an amino acid sequence having sequence identity to SEQ ID NO: 11636, wherein said polypeptide includes an amino acid sequence selected from the group consisting of AVSAC (SEQ ID NO: 11637) and AVGAC (SEQ ID NO: 11638). The invention includes a fragment of a polypeptide comprising SEQ ID NO: 11636, wherein said fragment includes an amino acid seuence selected from the group consisting of AVSAC (SEQ ID NO: 11637) and AVGAC (SEQ ID NO: 11638).

The invention includes a polynucleotide SEQ ID NO: 11639, or a fragment thereof or a sequence having sequence identity thereto. The invention includes a polypeptide encoded by the polynucleotide sequence set forth in SEQ ID NO: 11639, or a fragment thereof or a polypeptide sequence having sequence identity thereto.

The invention includes a polynucleotide set forth in SEQ ID NO: 11640, or a fragment thereof or a sequence having sequence identity thereto. The invention includes a polypeptide encoded by the polynucleotide sequence set forth in SEQ ID NO: 11640, or a fragment thereof or a polypeptide sequence having sequence identity thereto.

The invention includes each of the polynucleotides identified above. The invention includes each of the polynucleotides set forth in the sequence listing. The invention further

includes polynucleotides having sequence identity to each of the polynucleotides identified above. The degree of sequence identity is preferably greater than 50% (e.g., 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more).

The invention includes polynucleotide sequences comprising fragments of each of the polynucleotide sequences identified above. The fragments should comprise at least n consecutive polynucleotides from a particular SEQ ID NO:, and, depending on the sequence, n is 7 or more (e.g., 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 45, 50, 55, 60, 65, 70, 80, 90, 100, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300 or more).

The invention includes each of the amino acid sequences encoded by each of the polynucleotide sequences identified above. The invention includes each of the amino acid sequences encoded by each of the polynucleotide sequences set forth in the sequence listing. The invention further includes amino acid sequences having sequence identity to the amino acid sequences encoded by each of the polynucleotide sequences identified above. The degree of sequence identity is preferably greater than 50% (e.g., 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more). The invention further includes fragments of amino acid sequences encoded by

99% or more). The invention further includes fragments of amino acid sequences encoded by each of the polynucleotide sequences identified above. The fragments should comprise at least n consecutive amino acids from a particular SEQ ID NO:, and, depending on the sequence, n is 7 or more (e.g., 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 45, 50, 55, 60, 65, 70, 80, 90, 100, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300 or more).

The invention includes each of the amino acid sequences identified above. The invention includes each of the amino acid sequence set forth in the sequence listing. The invention further includes amino acid sequences having sequence identity to each of the amino acid sequences identified above. The degree of sequence identity is preferably greater than 50% (e.g., 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more).

The invention further includes fragments of the amino acid sequences identified above.

The fragments should comprise at least *n* consecutive amino acids from a particular SEQ ID

NO:, and, depending on the sequence, *n* is 7 or more (*e.g.*, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 45, 50, 55,

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60, 65, 70, 80, 90, 100, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300 or more).

The invention includes polynucleotides encoding each of the amino acid sequences identified above. The invention includes polynucleotides encoding each of the amino acid sequences set forth in the sequence listing. The invention further includes polynucleotides having sequence identity with each of the polynucleotides encoding each of the amino acid sequences identified above. The degree of sequence identity is preferably greater than 50% (e.g., 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more).

The invention further includes fragments of polynucleotides encoding each of the amino acid sequences identified above. The fragments should comprise at least n consecutive polynucleotides from a particular SEQ ID NO:, and, depending on the sequence, n is 7 or more (e.g., 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 45, 50, 55, 60, 65, 70, 80, 90, 100, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300 or more).

As described in more detail below, polynucleotides for use as primers and/or as probes may contain at least 4 or 8 contiguous nucleotides from a polynucleotide sequence of the invention e.g. at least 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, or 30 contiguous nucleotides and up to about 50, 75, 100, 200 contiguous nucleotides or more. While 6-8 nucleotides may be a workable length, sequences of 10-12 nucleotides are preferred, and about 13, 14, 15, 16, 17, 18, 19, 20, or 21 or more nucleotides or more appears optimal for hybridisation.

In one embodiment, the invention is directed to polynucleotides and amino acid sequences that do not consist entirely of a known SARS virus polynucleotide or amino acid sequence or of a known coronavirus polynucleotide or amino acid sequence. In one embodiment, the polynucleotides and amino acid sequences of the invention do not consist entirely of the sequence SEQ ID NO: 1. In another embodiment, the polynucleotides and amino acid sequences of the invention do not consist entirely of the sequence SEQ ID NO: 2. SEQ ID NO: 9967 is a SARS genome sequence of the Frankfurt (FRA) isolate (GenBank: AY310120). Compared to SEQ ID NO: 1, it differs at nucleotides 2546, 2590, 11437, 18954, 19073, 20585, 20899, 23209, 24922, 26589 & 28257; compared to SEQ ID NO:2, it differs at nucleotides 2560, 7922, 11451, 16625, 18968 & 19067. Further genome sequences have become available from GenBank, since this application was originally filed, under accession numbers including AY559097, AY559096, AY559095, AY559094, AY559093, AY559089, AY559088, AY559087, AY559086, AY559087, AY559088, AY559087, AY559086, AY559087, AY559081, AY274119,

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AY323977, AY291315, AY502932, AY502931, AY502930, AY502929, AY502928, AY502927, AY502926, AY502925, AY502924, AY502923, AY291451, AY390556, AY395003, AY395002, AY395001, AY395000, AY394999, AY394998, AY394997, AY394996, AY394995, AY394994, AY394993, AY394992, AY394991, AY394990, AY394989, AY394987, AY394986, AY394985, AY394983, AY394979, AY394978, AY508724, AY394850, AY463059, AY463060, AY313906, AY310120, AY461660, AY485278, AY485277, AY345988, AY345987, AY345986, AY282752, AY357076, AY357075, AY350750, AY304495, AY304488, AY304486, AY427439, AY283798, AY278491, AY278489, AY362699, AY362698, AY283797, AY283796, AY283795, AY283794, AY278741, AY351680, AP006561, AP006550, AP006559, AP006558, AP006557, AY278554, AY348314, AY338175, AY338174, AY321118, AY279354, AY278490, AY278487, AY297028, AY278488, and NC_004718.

In another embodiment, the invention is directed to polynucleotides that encode proteins which are not immunologically cross reactive with a protein of a mouse hepatitis virus, a bovine coronavirus or an avian infectious bronchitis virus. In another embodiment, the invention is directed to proteins which are not immunologically cross reactive with a protein of a mouse hepatitis virus, a bovine coronavirus or an avian infectious bronchitis virus.

Each of the polynucleotides identified above may be used to encode a portion of a fusion protein. Accordingly, the invention compries one or more of the polynucleotides identified above wherein the polynucleotides encoding for the start codon are removed. The invention further comprises one or more of the amino acids identified above wherein the starting methionine is removed.

Any of the polynucleotide or amino acid sequences discussed above may be used in vaccines for the treatment or prevention of SARS virus infection, including as a SARS viral antigen. Additionally, any of the polynucleotides or amino acid sequences discussed above may be used as diagnostic reagents, or in kits (comprising such reagents) or in methods used to diagnose or identify the presence or absence of a SARS virus in a biological sample.

SARS viral antigens of the invention may include a polypeptide with 99%, 95%, 90%, 85%, or 80% homology to one or more of the group consisting of the following proteins: nonstructural protein 2 (NS2); hemagglutinin-esterase glycoprotein (HE) (also referred to as E3), spike glycoprotein (S) (also referred to as E2), nonstructural region 4 (NS4), envelope (small membrane) protein (E) (also referred to as sM), membrane glycoprotein (M) (also referred to as E1), nucleocapsid phosphoprotein (N) or RNA dependent RNA polymerase (pol).

A detailed discussion of Coroavirus biology can be found in Fields Virology (2nd ed), Fields et al. (eds.), B.N. Raven Press, New York, NY., Chapter 35.

Another example of a SARS virus isolate is set forth in Example 1 below. The invention includes each of the polypeptide and polynucleotide sequences identified in Example 1. In

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addition, the invention includes vaccine formulations comprising one or more of the polypeptide or polynucleotide sequences identified in Example 1. The invention includes diagnostic regaents, kits (comprising such reagents) and methods which can be used to diagnose or identify the presence or absence of a SARS virus in a biological sample using one or more of the polypeptide or polynucleotide sequences identified in Example 1. The invention includes methods for the treatment or prevention of SARS virus infection utilizing small molecule viral inhibitors and combinations of small molecule viral inhibitors and kits for the treatment of SARS. The small molecule inhibitors may specifically target one or more of the polypeptides or polynucleotides identified in Example 1.

Further discussion of terms used in the application follows below.

"Respiratory Virus" as used herein refers to a virus capable of infecting the human respiratory tract. Respiratory Viral Antigens suitable for use in the invention include Severe Acute Respiratory Syndrome virus, coronavirus, influenza virus, human rhinovirus (HRV), parainfluenza virus (PIV), respiratory syncytial virus (RSV), adenovirus, metapneumovirus, and rhinovirus.

The terms "polypeptide", "protein" and "amino acid sequence" as used herein generally refer to a polymer of amino acid residues and are not limited to a minimum length of the product. Thus, peptides, oligopeptides, dimers, mulimers, and the like, are included within the definition. Both full-length proteins and fragments thereof are encompassed by the definition. Minimum fragments of polypeptides useful in the invention can be at least 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, or even 15 amino acids. Typically, polypeptides useful in this invention can have a maximum length suitable for the intended application. Generally, the maximum length is not critical and can easily be selected by one skilled in the art.

Polypeptides of the invention can be prepared in many ways e.g. by chemical synthesis (at least in part), by digesting longer polypeptides using proteases, by translation from RNA, by purification from cell culture (e.g. from recombinant expression), from the organism itself (e.g. after viral culture, or direct from patients), from a cell line source etc. A preferred method for production of peptides <40 amino acids long involves in vitro chemical synthesis (Bodanszky (1993) Principles of Peptide Synthesis (ISBN: 0387564314); Fields et al. (1997) Methods in Enzymology 289: Solid-Phase Peptide Synthesis. ISBN: 0121821900). Solid-phase peptide synthesis is particularly preferred, such as methods based on t-Boc or Fmoc (Chan & White (2000) Fmoc Solid Phase Peptide Synthesis ISBN: 0199637245) chemistry. Enzymatic synthesis (Kullmann (1987) Enzymatic Peptide Synthesis. ISBN: 0849368413) may also be used in part or in full. As an alternative to chemical synthesis, biological synthesis may be used e.g. the polypeptides may be produced by translation. This may be carried out in vitro or in vivo. Biological methods are in general restricted to the production of polypeptides based on L-amino

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acids, but manipulation of translation machinery (e.g. of aminoacyl tRNA molecules) can be used to allow the introduction of D-amino acids (or of other non natural amino acids, such as iodotyrosine or methylphenylalanine, azidohomoalanine, etc.) (Ibba (1996) Biotechnol Genet Eng Rev 13:197-216.). Where D-amino acids are included, however, it is preferred to use chemical synthesis. Polypeptides of the invention may have covalent modifications at the C-terminus and/or N-terminus, particularly where they are for in vivo administration e.g by attachment of acetyl or carboxamide, as in the Fuzeon TM product.

Reference to polypeptides and the like also includes derivatives of the amino acid sequences of the invention. Such derivatives can include postexpression modifications of the polypeptide, for example, glycosylation, acetylation, phosphorylation, and the like. Amino acid derivatives can also include modifications to the native sequence, such as deletions, additions and substitutions (generally conservative in nature), so long as the protein maintains the desired activity. These modifications may be deliberate, as through site-directed mutagenesis, or may be accidental, such as through mutations of hosts which produce the proteins or errors due to PCR amplification. Furthermore, modifications may be made that have one or more of the following effects: reducing toxicity; facilitating cell processing (e.g., secretion, antigen presentation, etc.); and facilitating presentation to B-cells and/or T-cells.

"Fragment" or "Portion" as used herein refers to a polypeptide consisting of only a part of the intact full-length polypeptide sequence and structure as found in nature. For instance, a fragment can include a C-terminal deletion and/or an N-terminal deletion of a protein.

A "recombinant" protein is a protein which has been prepared by recombinant DNA techniques as described herein. In general, the gene of interest is cloned and then expressed in transformed organisms, as described further below. The host organism expressed the foreign gene to produce the protein under expression conditions.

The term "polynucleotide", as known in the art, generally refers to a nucleic acid molecule. A "polynucleotide" can include both double- and single-stranded sequences and refers to, but is not limited to, cDNA from viral, prokaryotic or eukaryotic mRNA, genomic RNA and DNA sequences from viral (e.g. RNA and DNA viruses and retroviruses) or prokaryotic DNA, and especially synthetic DNA sequences. The term also captures sequences that include any of the known base analogs of DNA and RNA, and includes modifications such as deletions, additions and substitutions (generally conservative in nature), to the native sequence, so long as the nucleic acid molecule encodes a therapeutic or antigenic protein. These modifications may be deliberate, as through site-directed mutagenesis, or may be accidental, such as through mutations of hosts that produce the antigens. Modifications of polynucleotides may have any number of effects including, for example, facilitating expression of the polypeptide product in a host cell.

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Polynucleotides of the invention may be prepared in many ways e.g. by chemical synthesis (e.g. phosphoramidite synthesis of DNA) in whole or in part, by digesting longer nucleic acids using nucleases (e.g. restriction enzymes), by joining shorter nucleic acids or nucleotides (e.g. using ligases or polymerases), from genomic or cDNA libraries, etc.

A polynucleotide can encode a biologically active (e.g., immunogenic or therapeutic) protein or polypeptide. Depending on the nature of the polypeptide encoded by the polynucleotide, a polynucleotide can include as little as 10 nucleotides, e.g., where the polynucleotide encodes an antigen.

By "isolated" is meant, when referring to a polynucleotide or a polypeptide, that the indicated molecule is separate and discrete from the whole organism with which the molecule is found in nature or, when the polynucleotide or polypeptide is not found in nature, is sufficiently free of other biological macromolecules so that the polynucleotide or polypeptide can be used for its intended purpose. The polynucleotides and polypeptides of the invention are preferably isolated polynucleotides and isolated polypeptides.

"Antibody" as known in the art includes one or more biological moieties that, through chemical or physical means, can bind to or associate with an epitope of a polypeptide of interest. The antibodies of the invention include antibodies which specifically bind to a SARS viral antigen. The term "antibody" includes antibodies obtained from both polyclonal and monoclonal preparations, as well as the following: hybrid (chimeric) antibody molecules (see, for example, Winter et al. (1991) Nature 349: 293-299; and US Patent No. 4,816,567; F(ab')2 and F(ab) fragments; Fv molecules (non-covalent heterodimers, see, for example, Inbar et al. (1972) Proc Natl Acad Sci USA 69:2659-2662; and Ehrlich et al. (1980) Biochem 19:4091-4096); singlechain Fv molecules (sFv) (see, for example, Huston et al. (1988) Proc Natl Acad Sci USA 85:5897-5883); dimeric and trimeric antibody fragment constructs; minibodies (see, e.g., Pack et al. (1992) Biochem 31:1579-1584; Cumber et al. (1992) J Immunology 149B: 120-126); humanized antibody molecules (see, for example, Riechmann et al. (1988) Nature 332:323-327; Verhoeyan et al. (1988) Science 239:1534-1536; and U.K. Patent Publication No. GB 2,276,169. published 21 September 1994); and, any functional fragments obtained from such molecules, wherein such fragments retain immunological binding properties of the parent antibody molecule. The term "antibody" further includes antibodies obtained through non-conventional processes, such as phage display.

As used herein, the term "monoclonal antibody" refers to an antibody composition having a homogeneous antibody population. The term is not limited regarding the species or source of the antibody, nor is it intended to be limited by the manner in which it is made. Thus, the term encompasses antibodies obtained from murine hybridomas, as well as human monoclonal

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antibodies obtained using human rather than murine hybridomas. See, e.g., Cote, et al. Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, 1985, p 77.

An "immunogenic composition" as used herein refers to a composition that comprises an antigenic molecule where administration of the composition to a subject results in the development in the subject of a humoral and/or a cellular immune response to the antigenic molecule of interest. The immunogenic composition can be introduced directly into a recipient subject, such as by injection, inhalation, oral, intranasal or any other parenteral, mucosal or transdermal (e.g., intra-rectally or intra-vaginally) route of administration.

The term "derived from" is used to identify the source of molecule (e.g., a molecule can be derived from a polynucleotide, polypeptide, an immortalized cell line can be derived from any tissue, etc.). A first polynucleotide is "derived from" a second polynucleotide if it has the same or substantially the same basepair sequence as a region of the second polynucleotide, its cDNA, complements thereof, or if it displays sequence identity as described above. Thus, a first polynucleotide sequence is "derived from" a second sequence if it has (i) the same or substantially the same sequence as the second sequence or (ii) displays sequence identity to polypeptides of that sequence.

A first polypeptide is "derived from" a second polypeptide if it is (i) encoded by a first polynucleotide derived from a second polynucleotide, or (ii) displays sequence identity to the second polypeptides as described above. Thus, a polypeptide (protein) is "derived from" a particular SARS virus if it is (i) encoded by an open reading frame of a polynucleotide of that SARS virus, or (ii) displays sequence identity, as described above, to polypeptides of that SARS virus.

Both polynucleotide and polypeptide molecules can be physically derived from a SARS virus or produced recombinantly or synthetically, for example, based on known sequences.

A cultured cell or cell line is "derived from" another cell, cells or tissue if it is originally obtained from existing cells or tissue. Non-limiting examples of tissue that cells may be derived from include skin, retina, liver, kidney, heart, brain, muscle, intestinal, ovary, breast, prostate, cancerous tissue, tissue infected with one or more pathogens (e.g., viruses, bacteria etc.) and the like. The cells described herein may also be derived from other cells including, but not limited to, primary cultures, existing immortalized cells line and/or other isolated cells.

An "antigen" refers to a molecule containing one or more epitopes (either linear, conformational or both) that will stimulate a host's immune system to make a humoral and/or cellular antigen-specific response. The term is used interchangeably with the term "immunogen." Normally, an epitope will include between about 3-15, generally about 5-15 amino acids. A B-cell epitope is normally about 5 amino acids but can be as small as 3-4 amino acids. A T-cell epitope, such as a CTL epitope, will include at least about 7-9 amino acids, and a

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helper T-cell epitope at least about 12-20 amino acids. Normally, an epitope will include between about 7 and 15 amino acids, such as, 9, 10, 12 or 15 amino acids. The term "antigen" denotes both subunit antigens, (i.e., antigens which are separate and discrete from a whole organism with which the antigen is associated in nature), as well as, killed, attenuated or inactivated bacteria, viruses, fungi, parasites or other microbes as well as tumor antigens, including extracellular domains of cell surface receptors and intracellular portions that may contain T-cell epitopes. Antibodies such as anti-idiotype antibodies, or fragments thereof, and synthetic peptide mimotopes, which can mimic an antigen or antigenic determinant, are also captured under the definition of antigen as used herein. Similarly, an oligonucleotide or polynucleotide that expresses an antigen or antigenic determinant in vivo, such as in gene therapy and DNA immunization applications, is also included in the definition of antigen herein.

An "immunological response" to an antigen or composition is the development in a subject of a humoral and/or a cellular immune response to an antigen present in the composition of interest. For purposes of the present invention, a "humoral immune response" refers to an immune response mediated by antibody molecules, including secretory (IgA) or IgG molecules, while a "cellular immune response" is one mediated by T-lymphocytes and/or other white blood cells. One important aspect of cellular immunity involves an antigen-specific response by cytolytic T-cells ("CTL"s). CTLs have specificity for peptide antigens that are presented in association with proteins encoded by the major histocompatibility complex (MHC) and expressed on the surfaces of cells. CTLs help induce and promote the destruction of intracellular microbes, or the lysis of cells infected with such microbes. Another aspect of cellular immunity involves an antigen-specific response by helper T-cells. Helper T-cells act to help stimulate the function, and focus the activity of, nonspecific effector cells against cells displaying peptide antigens in association with MHC molecules on their surface. A "cellular immune response" also refers to the production of cytokines, chemokines and other such molecules produced by activated T-cells and/or other white blood cells, including those derived from CD4+ and CD8+ T-cells. In addition, a chemokine response may be induced by various white blood or endothelial cells in response to an administered antigen.

II. VACCINE FORMULATIONS

The invention relates to vaccine formulations for the treatment or prevention of Severe Acute Respiratory Syndrome (SARS). Vaccine formulations of the invention include an inactivated (or killed) SARS virus, an attenuated SARS virus, a split SARS virus preparation and a recombinant or purified subunit formulation of one or more SARS viral antigens. The invention includes polypeptides and polynucleotides encoding for SARS viral antigens and

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immunogenic fragments thereof. Expression and delivery of the polynucleotides of the invention may be facilitated via viral vectors and/or viral particles, including Virus Like Particles (VLPs).

A. Inactivated (or Killed) SARS Vaccines

The invention includes a composition comprising an inactivated (or killed) SARS virus and methods for the production thereof. Inactivated SARS viral compositions can be used as prophylactic or therapeutic SARS virus vaccine. Preferably the inactivated SARS virus vaccine composition comprises an amount of inactivated SARS virus which, before inactivation, is equivalent to a virus titer of from about 4 to 7 logs plaque forming units (PFU) or 4 to 7 logs tissue culture infectious dose 50 (TCID₅₀) per milliliter. More preferably, before inactivation the virus titer is from 4 to 11, 7 to 11 or 9 to 11 PFU or TCID₅₀. Still more preferably the inactivated SARS virus vaccine composition comprises an amount of inactivated SARS virus which, before inactivation, is equivalent to a virus titer of from about 5 to 9 PFU or 5 to 9 TCID₅₀ per milliliter. In one embodiment, the PFU or TCID₅₀ per milliliter. Upon concentration of the viral harvest, the PFU or TCID₅₀ is preferably 8 to 11, still more preferably about 9 PFU or TCID₅₀ per milliliter. The vaccine composition comprises a sufficient amount of the SARS virus antigen to produce an immunological response in a primate.

Methods of inactivating or killing viruses are known in the art to destroy the ability of the viruses to infect mammalian cells. Such methods include both chemical or physical means. Chemical means for inactivating a SARS virus include treatment of the virus with an effective amount of one or more of the following agents: detergents, formaldehyde, formalin, β -propiolactone, or UV light. Additional chemical means for inactivation include treatment with methylene blue, psoralen, carboxyfullerene (C60) or a combination of any thereof. Other methods of viral inactivation are known in the art, such as for example binary ethylamine, acetyl ethyleneimine, or gamma irradiation.

For example formaldehyde may be used at concentrations such as 0.1 to 0.02%, preferably at 0.02 to 0.1%, and still more preferably at 0.04 to 0.05%. The inactivating agent is added to virus containing culture supernatants prior to or after harvesting said culture supernatants from vessels used for virus propagation, either with or without a step of cell disruption for release of cell-associated virus prior to harvesting. Further, the inactivating agent may be added after said culture supernatants have been stored frozen and thawed, or after one or more steps of purification to remove cell contaminants. Preferably, however, formaldehyde is added after removal of cells and cellular debris or after one or more purification steps. After addition of formaldehyde, the virus containing mixture is transferred into an incubation vessel and incubated at refrigeration temperatures (e.g. +2 to 8°C) or alternatively at elevated temperatures, such as ambient temperatures between approximately 20 and 30°C or at 33°C to 37°C for a period of 12 -105-

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hours to 7 days, whereby the temperature chosen should be adjusted to the duration of incubation. Prefered conditions are e.g. $+2-8^{\circ}$ C for 3-7 days (prefered are 3-7days), ambient temperatures and incubation for 16 hours to 3 days (prefered 24-48 hours), or 35-37°C for 12-36 hours. If it is desirable to remove excess formalin, sodium thiosulfate or sodium metabisulfite at equimolar or 1.5-fold molar concentration (relative to formaldehyde) may be added after completing the inactivation process.

For example, β -propiolactone may be used at concentrations such as 0.01 to 0.5%, preferably at 0.5% to 0.2%, and still more preferably at 0.025 to 0.1%. The inactivating agent is added to virus containing culture supernatants (virus material) prior to or after harvesting said culture supernatants from vessels used for virus propagation, either with or without a step of cell disruption for release of cell-associated virus prior to harvesting. Further, the inactivating agent may be added after said culture supernatants have been stored frozen and thawed, or after one or more steps of purification to remove cell contaminants. β -propiolactone is added to the virus material, with the adverse shift in pH to acidity being controlled with sodium hydroxide (e.g., 1 N NaOH), a Tris-buffer or sodium bicarbonate solution. After transfering the mixture to another inactivation vessel, the combined inactivating agent-virus materials are incubated at temperatures from 4°C to 37°C, for incubation times of preferably 24 to 72 hours.

Another inactivant which may be used is binary ethyleneimine (BEI). Equal volumes of a 0.2 molar bromoethylamine hydrobromide solution and a 0.4 molar sodium hydroxide solution are mixed and incubated at about 37°C. for 60 minutes. The resulting cyclized inactivant is binary ethyleneimine, which is added to the virus materials at 0.5 to 4 percent, and preferably at 1 to 3 percent, volume to volume. The inactivating virus materials are held from about 4°C to 37°C for 24 to 72 hours with periodic agitation. At the end of this incubation 20 ml. of a sterile 1 molar sodium thiosulfate solution was added to insure neutralization of the BEI.

In one embodiment, the invention includes an inactivating method is designed to maximize exposure of the virus to the inactivating agent and to minimize long-term exposure of the temperature sensitive SARS virus particles to elevated temperatures. The invention includes an inactivation method comprising exposing the virus to the inactivation agent (such as BPL) for 12 to 24 hours at refrigeration temperatures followed by hydrolysis of any residual inactivating agent by elevating the temperature for only 3 hours. Preferably, the refrigeration temperatures are between 0 and 8°C, more preferably around 4°C. Preferably, the elevated temperature is between 33 and 41°C, more preferably around 37°C. As assessed by a test for residual infectious virus using 10 ml aliquots of the inactivated preparation, the method is able to inactivate SARS-CoV in raw cell culture harvests below a theoretical limit of 0.03 infectious units/ml.

Diluted and undiluted samples of the inactivated virus materials are added to susceptible cell (tissue) culture (e.g., VERO) to detect any non-inactivated virus. The cultured cells are

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passaged multiple times and examined for the presence of SARS virus based on any of a variety of methods, such as, for example, cytopathic effect (CPE) and antigen detection (e.g., via fluoroscent antibody conjugates specific for SARS virus). Such tests allow determination of complete virus inactivation.

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Prior to inactivation, the SARS virus will be cultured in a mammalian cell culture. The cell culture may be adherently growing cells or cells growing in suspension. Preferably the cells are of mammalian origin, but may also be derived from avian (e.g., hens' cells such as hens' embryo cells (CEF cells)), amphibian, reptile, insect, or fish sources. Mammalian sources of cells include, but are not limited to, human or non-human primate (e.g., MRC-5 (ATCC CCL-171), WI-38 (ATCC CCL-75), HeLa cells, human diploid cells, fetal rhesus lung cells (e.g. ATCC CL-160), human embryonic kidney cells (293 cells, typically transformed by shead adenovirus type 5 DNA), VERO cells (e.g., from monkey kidneys), horse, cow (e.g., MDBK cells), sheep, dog (e.g., MDCK cells from dog kidneys, ATCC CCL-34 MDCK (NBL2) or MDCK 33016, deposit number DSM ACC 2219 as described in WO 97/37001), cat, and rodent (e.g., hamster cells such as BHK21-F, HKCC cells, or Chinese hamster ovary cells (CHO cells)), and may be obtained from a wide variety of developmental stages, including for example, adult, neonatal, fetal, and embryo.

In certain embodiments the cells are immortalized (e.g., PERC.6 cells are described, for example, in WO 01/38362 and WO 02/40665, incorporated by reference herein in their entireties, as well as deposited under ECACC deposit number 96022940), or any other cell type immortalized using the techniques described herein.

In preferred embodiments, mammalian cells are utilized, and may be selected from and/or derived from one or more of the following non-limiting cell types: fibroblast cells (e.g., dermal, lung), endothelial cells (e.g., aortic, coronary, pulmonary, vascular, dermal microvascular, umbilical), hepatocytes, keratinocytes, immune cells (e.g., T cell, B cell, macrophage, NK, dendritic), mammary cells (e.g., epithelial), smooth muscle cells (e.g., vascular, aortic, coronary, arterial, uterine, bronchial, cervical, retinal pericytes), melanocytes, neural cells (e.g., astrocytes), prostate cells (e.g., epithelial, smooth muscle), renal cells (e.g., epithelial, mesangial, proximal tubule), skeletal cells (e.g., chondrocyte, osteoclast, osteoblast), muscle cells (e.g., myoblast, skeletal, smooth, bronchial), liver cells, retinoblasts, and stromal cells. WO 97/37000 and WO 97/37001, incorporated by reference herein in their entireties, describe production of animal cells and cell lines that capable of growth in suspension and in serum free media and are useful in the production and replication of viruses.

Preferably, the SARS viruses of the invention are grown on VERO cells or fetal rhesus kidney cells.

Culture conditions for the above cell types are well-described in a variety of publications, or alternatively culture medium, supplements, and conditions may be purchased commercially, such as for example, as described in the catalog and additional literature of Cambrex Bioproducts (East Rutherford, NJ).

In certain embodiments, the host cells used in the methods described herein are cultured in serum free and/or protein free media. A medium is referred to as a serum-free medium in the context of the present invention in which there are no additives from serum of human or animal origin. Protein-free is understood to mean cultures in which multiplication of the cells occurs with exclusion of proteins, growth factors, other protein additives and non-serum proteins. The cells growing in such cultures naturally contain proteins themselves.

Known serum-free media include Iscove's medium, Ultra-CHO medium (BioWhittaker) or EX-CELL (JRH Bioscience). Ordinary serum-containing media include Eagle's Basal Medium (BME) or Minimum Essential Medium (MEM) (Eagle, Science, 130, 432 (1959)) or Dulbecco's Modified Eagle Medium (DMEM or EDM), which are ordinarily used with up to 10% fetal calf serum or similar additives. Optionally, Minimum Essential Medium (MEM) (Eagle, Science, 130, 432 (1959)) or Dulbecco's Modified Eagle Medium (DMEM or EDM) may be used without any serum containing supplement. Protein-free media like PF-CHO (JHR Bioscience), chemically-defined media like ProCHO 4CDM (BioWhittaker) or SMIF 7 (Gibco/BRL Life Technologies) and mitogenic peptides like Primactone, Pepticase or HyPepTM (all from Quest International) or lactalbumin hydrolyzate (Gibco and other manufacturers) are also adequately known in the prior art. The media additives based on plant hydrolyzates have the special advantage that contamination with viruses, mycoplasma or unknown infectious agents can be ruled out.

The cell culture conditions to be used for the desired application (temperature, cell density, pH value, etc.) are variable over a very wide range owing to the suitability of the cell line employed according to the invention and can be adapted to the requirements of the SARS virus.

The method for propagating the SARS virus in cultured cells (e.g., mammalian cells) includes the steps of inoculating the cultured cells with SARS virus, cultivating the infected cells for a desired time period for virus propagation, such as for example as determined by SARS virus titer or SARS virus antigen expression (e.g., between 24 and 168 hours after inoculation) and collecting the propagated virus. The cultured cells are inoculated with a SARS virus (measured by PFU or TCID₅₀) to cell ratio of 1:10000 to 1:10. A lower range of ratios may also be used e.g. 1:500 to 1:1, preferably 1:100 to 1:5, more preferably 1:50 to 1:10. The SARS virus is added to a suspension of the cells or is applied to a monolayer of the cells, and the virus is absorbed on the cells for at least 60 minutes but usually less than 300 minutes, preferably between 90 and 240 minutes at 25°C to 40°C, more preferably 28°C to 37°C, still more

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preferably at about 33 °C. The infected cell culture (e.g., monolayers) may be treated either by freeze-thawing or by enzymatic action to increase the viral content of the harvested culture supernatants. The harvested fluids are then either inactivated or stored frozen.

A comparison of SARS infected Vero cells grown with and without fetal calf serum ("FCS") is shown in FIGURE 26A. Briefly, Vero cells were split the day before infection and cultivated in T175 flasks. Infection of 90% confluent Vero cell monolayers the following day was performed with a SARS-CoV seed stock (strain FRA, passage 4, Accession number AY310120), with or without 3% FCS (Fig. 26A). The addition of FCS to the cell media showed little impact on virus yield.

Cultured cells may be infected at a multiplicity of infection ("m.o.i.") of about 0.0001 to 10, preferably 0.002 to 5, more preferably to 0.001 to 2. Still more preferably, the cells are infected at a m.o.i of about 0.01. A comparison of viral yield at varying m.o.i. levels is shown in FIGURE 26B.

Infected cells may be harvested 30 to 60 hours post infection. Preferably, the cells are harvested 34-48 hours post infection. Still more preferably, the cells are harvested 38 to 40 hours post infection. See FIGURE 26C.

Methods of purification of inactivated virus are known in the art and may include one or more of, for instance gradient centrifugation, ultracentrifugation, continuous-flow ultracentrifugation and chromatography, such as ion exchange chromatography, size exclusion chromatography, and liquid affinity chromatography. Additional method of purification include ultrafiltration and dialfiltration. See JP Gregersen "Herstellung von Virussimpfstoffen aus Zellkulturen" Chapter 4.2 in Pharmazeutische Bioteonologie (eds. O. Kayser and RH Mueller). Wissenschaftliche Verlagsgesellschaft, Stuttgart, 2000. See also, O'Neil et al., "Virus Harvesting and Affinity Based Liquid Chromatography. A Method for Virus Concentration and Purification", Biotechnology (1993) 11:173-177; Prior et al., "Process Development for Manufacture of Inactivated HIV-1", Pharmaceutical Technology (1995) 30-52; and Majhdi et al., "Isolation and Characterization of a Coronavirus from Elk Calves with diarrhea" Journal of Clinical Microbiology (1995) 35(11): 2937-2942.

Other examples of purification methods suitable for use in the invention include polyethylene glycol or ammonium sulface precipitation (see Trepanier et al., "Concentration of human respiratory syncytial virus using ammonium sulfate, polyethylene glycol or hollow fiber ultrafiltration" Journal of Virological Methods (1981) 3(4):201-211; Hagen et al., "Optimization of Poly(ethylene glycol) Precipitation of Hepatitis Virus Used to prepare VAQTA, a Highly Purified Inactivated Vaccine" Biotechnology Progress (1996) 12:406-412; and Carlsson et al., "Purification of Infectious Pancreatic Necrosis Virus by Anion Exchange Chromatography Increases the Specific Infectivity" Journal of Virological Methods (1994) 47:27-36) as well as

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ultrafiltration and microfiltration (see Pay et al., Developments in Biological Standardization (1985) 60:171-174; Tsurumi et al., "Structure and filtration performances of improved cuprammonium regenerated cellulose hollow fibre (improved BMM hollow fibre) for virus removal" Polymer Journal (1990) 22(12):1085-1100; and Makino et al., "Concentration of live retrovirus with a regenerated cellulose hollow fibre, BMM", Archives of Virology (1994) 139(1-2):87-96.).

Preferably, the virus is purified using chromatography, such as ion exchange chromatography. Chromatic purification allows for the production of large volumes of virus containing suspension. The viral product of interest can interact with the chromatic medium by a simple adsorption/desorption mechanism, and large volumes of sample can be processed in a single load. Contaminants which do not have affinity for the adsorbent pass through the column. The virus material can then be eluted in concentrated form.

Preferred anion exchange resins for use in the invention include DEAE, EMD TMAE. Preferred cation exchange resins may comprise a sulfonic acid-modified surface. In one embodiment, the virus is purified using ion exchange chromatography comprising a strong anion exchange resin (e.g. EMD TMAE) for the first step and EMD-SO₃ (cation exchange resin) for the second step. A metal-binding affinity chromatography step can optionally be included for further purification. (See, e.g., WO 97/06243).

A preferred resin for use in the invention is FractogelTM EMD. This synthetic methacrylate based resin has long, linear polymer chains (so-called "tentacles") covalently attached. This "tentacle chemistry" allows for a large amount of sterically accessible ligands for the binding of biomolecules without any steric hindrance. This resin also has improved pressure stability.

Column-based liquid affinity chromatography is another preferred purification method for use in the invention. One example of a resin for use in this purification method is MatrexTM CellufineTM Sulfate (MCS). MCS consists of a rigid spherical (approx. 45-105 μ m diameter) cellulose matrix of 3,000 Dalton exclusion limit (its pore structure excludes macromolecules), with a low concentration of sulfate ester functionality on the 6-position of cellulose. As the functional ligand (sulfate ester) is relatively highly dispersed, it presents insufficient cationic charge density to allow for most soluble proteins to adsorb onto the bead surface. Therefore the bulk of the protein found in typical virus pools (cell culture supernatants, e.g. pyrogens and most contaminating proteins, as well as nucleic acids and endotoxins) are washed from the column and a degree of purification of the bound virus is achieved.

The rigid, high-strength beads of MCS tend to resist compression. The pressure/flow characteristics the MCS resin permit high linear flow rates allowing high-speed processing, even in large columns, making it an easily scalable unit operation. In addition a chromatographic purification step with MCS provides increased assurance of safety and product sterility, avoiding

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excessive product handling and safety concerns. As endotoxins do not bind to it, the MCS purification step allows a rapid and contaminant free depyrogenation. Gentle binding and elution conditions provide high capacity and product yield. The MCS resin therefore represents a simple, rapid, effective, and cost-saving means for concentration, purification and depyrogenation. In addition, MCS resins can be reused repeatedly.

The inactivated virus may be further purified by gradient centrifugation, preferably density gradient centrifugation. For commercial scale operation a continuous flow sucrose gradient centrifugation would be the preferred option. This method is widely used to purify antiviral vaccines and is known to the expert in the field (See JP Gregersen "Herstellung von Virussimpfstoffen aus Zellkulturen" Chapter 4.2 in Pharmazeutische Biotecnologie (eds. O. Kayser and RH Mueller) Wissenschaftliche Verlagsgesellschaft, Stuttgart, 2000.)

The density gradient centrifugation step may be performed using laboratory or commercial scale gradient centrifugation equipment. For example, a swinging bucket rotor, a fixed angle rotor, or a vertical tube rotor, particularly for laboratory scale production of the virus. Preferably, the gradient centrifugation step is performed using a swinging bucket rotor. This type of rotor has a sufficiently long pathlength to provide high quality separations, particularly with multicomponent samples. In addition, swinging bucket rotors have greatly reduced wall effects, and the contents do not reorient during acceleration and deceleration. Because of their longer pathlength, separations take longer compared to fixed angle or vertical tube rotors. The prepared sucrose solutions are controlled via refractometer on their sucrose concentration.

Sucrose gradients for density gradient centrifugation, such as in a swinging bucket centrifuge tubes may be formed prior to centrifugation by the use of a gradient former (continuous/linear). The volume of sample which can be applied to the gradient in a swinging bucket rotor tube is a function of the cross-sectional area of the gradient that is exposed to the sample. If the sample volume is too high, there is not sufficient radial distance in the centrifuge tube for effective separation of components in a multicomponent sample.

An approximate sample volume for swinging bucket rotor SW 28 is 1-5 ml per tube (with a tube diameter of 2.54 cm). The sample is applied to the gradient by pipetting the volume on top of the gradient. The blunt end of the pipette is placed at 45-60° angle to the tube wall, approximately 2-3 mm above the gradient. The sample is injected slowly and allowed to run down the wall of the tube onto the gradient. After centrifugation gradient fractions are recovered by carefully inserting a gauge needle until the bottom of the tube and starting to collect fractions of 2 ml by pumping the liquid from the tube into falcon tubes.

Sucrose density gradients suitable for use with this density gradient centrifugation purification step include 0-60%, 5-60%, 15-60%, 0-50%, 5-50%, 15-50%, 0-40%, 5-40%, and 15-40%. Preferably, the sucrose density gradient is 15-40%, 5-40% or 0-40%.

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Alternatively, a discontinuous sucrose density gradient may be used for purification. A discontinuous sucrose density scheme provides for discrete, overlaying layers of differing sucrose concentrations. In one example, a first layer of 50% sucrose is covered by a second layer of 40% sucrose; the second layer is covered by a third layer of 20% sucrose; the third layer is covered by a fourth layer of 10% sucrose; and the fourth layer is covered by the solution containing the virus to be purified.

In one embodiment, inactivated virus is purified by a method comprising a first step of chromatography purification and a second step of gradient centrifugation. Preferably the first step comprises liquid affinity chromatography, such as MCS. Preferably, the second step comprises density gradient centrifugation using a swinging bucket rotor.

Additional purification methods which may be used to purify inactivated SARS virus include the use of a nucleic acid degrading agent, preferably a nucleic acid degrading enzyme, such as a nuclease having DNase and RNase activity, or an endonuclease, such as from Serratia marcescens, commercially available as BenzonaseTM, membrane adsorbers with anionic functional groups (e.g. SartobindTM) or additional chromatographic steps with anionic functional groups (e.g. DEAE or TMAE). An ultrafiltration/dialfiltration and final sterile filtration step could also be added to the purification method.

Preferably, the purification includes treatment of the SARS viral isolate with one or more nucleic acid degrading enzymes. These enzymes may be used to reduce the level of host cell nucleic acid in the viral purification process. Nucleic acid digesting enzymes for use in cell culture are known in the art and include, for example, BenzonaseTM.

The treatment of the virus with the nucleic acid degrading enzyme and inactivating agent can be performed by a sequential treatment or in a combined or simultaneous manner.

Preferably, the nucleic acid degrading agent is added to the virus preparation prior to the addition of the inactivating agent.

The purified viral preparation of the invention is substantially free of contaminating proteins derived from the cells or cell culture and preferably comprises less than about 1000, 500, 250, 150, 100, or 50 pg cellular nucleic acid / µg virus antigen, preferably less than about 1000, 500, 250, 150, 100, or 50 pg cellular nucleic acid/ dose. Still more preferably, the purified viral preparation comprises less than about 20 pg, and even more preferably, less than about 10 pg. Methods of measuring host cell nucleic acid levels in a viral sample are known in the art. Standardized methods approved or recommended by regulatory authorities such as the WHO or the FDA are preferred.

The invention includes an inactivated vaccine composition comprising a prophylactically effective amount of SARS viral antigen, preferably spike or an immunogenic fragment thereof. The SARS viral antigen is preferably present in a concentration amount of 0.1 to 50 μ g

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antigen/dose, more preferably 0.3 to 30 μg antigen/dose. Still more preferably, the antigen is about 15 μg /dose.

In one embodiment, a lower concentration of SARS viral antigen is used in inactivated vaccine compositions of the invention. Such lower concentration vaccines may optionally comprise an adjuvant to boost the host immune response to the antigen. In such a "low dose" vaccine, the SARS viral antigen is preferably present in a concentration of less than $15 \,\mu g$ antigen/dose, (i.e., less than 10, 7.5, 5 or $3 \,\mu g$ antigen/dose.

The inactivated vaccine preparations of the invention may further comprise a stabilizer to preserve the integrity of the immunogenic proteins in the inactivated viral preparation. Stabilizers suitable for use in vaccines are known in the art and may include, for example, buffers, sugars, sugar alcohols, and amino acids. Stabilizing buffers are preferably adjusted to a physiological pH range and may include phosphate buffers, Tris buffers, TE (Tris/EDTA), TEN (Tris/NaCl/EDTA) and Earle's salt solution. Stabilizing sugars may include, for example, one or more of saccharose, glucose, fructose, dextranes, dextranesulphate, and trehalose. Stabilizing sugar alcohols may include, for example, Xylite/Xylitole, Mannite/Mannitol, Sorbite/Sorbitol, and Glycerol. Amino acids suitable for use in the invention include, for example, L-glutamine, arginine, cysteine, and lysine. Additional stabilizers which may be used in the invention include Tartaric acid, Pluronic F 68, and Tween 80.

SARS viral isolates which may be used for the inactivated viral preparations of the invention may be obtained and identified by any of the mechanisms described supra. For example, a SARS isolate may be obtained from a clinical sample and plaque purified. Such methods of viral isolation are known in the art.

Further purification procedures can be applied to ensure the seed virus used for preparation of the vaccine does not contain, for example, unwanted adventitious agents. In one embodiment, viral RNA from the viral isolate can be isolated from the virus, purified (and, optionally, the sequence verified through PCR or other means) and then introduced into a suitable cell culture.

As an example of this technique, a clinical viral sample is plaque purified and amplified on vero cells to generate a sufficient amount of the viral sample for analysis. Cellular remnants are then cleared from the supernatant by centrifugation. The virus can then be pelleted by ultracentrifugation and the pellet resuspended in PBS. After further centrifugation purification, the virus containing fraction is treated with a DNase (and optionally also an RNase). Viral RNA is then isolated from this fraction and transfected into a host cell.

Examples 2 and 3 provide an illustration of purification of inactivated whole SARS virus using MCS chromatography resin purification followed by density gradient ultracentrifugation.

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Routes and methods of immunization of the vaccines of the invention are discussed in more detail in a section below. Examples 4 and 5 provide illustrations of a mouse immunization scheme with the inactivated SARS virus of the invention.

B. Attenuated SARS Vaccines

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The invention includes a composition comprising an attenuated SARS virus. This composition can be used as a prophylactic or therapeutic SARS virus vaccine. Methods of attenuating viruses are known in the art. Such methods include serial passage of the SARS virus in cultured cells (e.g., mammalian cell culture, preferably fetal rhesus kidney cells or VERO cells-see the discussion in Section A above regarding culture of SARS virus), until the SARS virus demonstrates attenuated function. The temperature at which the virus is grown can be any temperature at which with tissue culture passage attenuation occurs. Attenuated function of the SARS virus after one or more passages in cell culture can be measured by one skilled in the art. As used herein, attenuation refers to the decreased virulence of the SARS virus in a human subject. Evidence of attenuated function may be indicated by decreased levels of viral replication or by decreased virulence in an animal model.

Other methods of producing an attenuated SARS virus include passage of the virus in cell culture at sub-optimal or "cold" temperatures and introduction of attenuating mutations into the SARS viral genome by random mutagenesis (e.g., chemical mutagenesis) or site specific directed mutagenesis. Preparation and generation of attenuated RSV vaccines (the methods of which will generally applicable to SARS virus) are disclosed in, for example, EP 0 640 128, US Patent No. 6,284,254, US Patent No. 5,922,326, US Patent No. 5,882,651.

The attenuated derivatives of SARS virus are produced in several ways, such as for example, by introduction of temperature sensitive-mutations either with or without chemical mutagenesis (e.g., 5-fluorouracil), by passage in culture at "cold" temperatures. Such cold adaptation includes passage at temperatures between about 20°C to about 32°C, and preferably between temperatures of about 22°C to about 30°C, and most preferably between temperatures of about 24°C and 28°C. The cold adaptation or attenuation may be performed by passage at increasingly reduced temperatures to introduce additional growth restriction mutations. The number of passages required to obtain safe, immunizing attenuated virus is dependent at least in part on the conditions employed. Periodic testing of the SARS virus culture for virulence and immunizing ability in animals (e.g., mouse, primate) can readily determine the parameters for a particular combination of tissue culture and temperature. The attenuated vaccine will typically be formulated in a dose of from about 10³ to 106 PFU or TCID50, or more for maximal efficacy.

Attenuated virus vaccines for SARS-CoV also are produced by creating virus chimeras comprising sequences derived from at least two different coronaviruses, one of which is a SARS-CoV. For example, a virus chimera is produced that comprises nonstructural protein encoding

genes derived from a first coronavirus (e.g., murine, bovine, porcine, canine, feline, avian coronavirus) and one or more structural protein encoding genes (e.g., spike, E, M) from a SARS-CoV. Alternatively, the virus chimera may comprise sequences derived from a human coronavirus that is not a SARS-CoV (e.g., OC43, 229E) together with sequences from a SARS-CoV. Chimeric coronaviruses of the present invention are generated by a variety of methods, including for example allowing for natural RNA recombination in a eukaryotic (e.g., mammalian) cell that contains RNA from each of the parental coronaviruses (e.g., following infection) or by using standard molecular biology techniques known to those of skill in the art to engineer desired virus chimeras (or portions thereof) as cDNA clones, which may then be used to produce infectious virus (see for example, US 6593111 B2; Yount et al., 2003, Proc. Natl. Acad. Sci. USA 100(22):12995-13000). An attenuated phenotype of the coronavirus chimeras described herein can be readily measured by one of skill in the art.

Attenuated viruses can be also generated by deleting one or more open reading frames (ORFs) that are not essential for viral replication. Preferably, these deletions occur in the structural region of the genome, such as ORF 3a, 3b, 6, 7a, 7b, 8a, 8b, 9b. See e.g., Haijema BJ, Volders H, Rottier PJ. J Virol. (2004) 78(8):3863-71; and de Haan, C. A., P. S. Masters, X. Shen, S. Weiss, and P. J. Rottier, "The group-specific murine coronavirus genes are not essential, but their deletion, by reverse genetics, is attenuating in the natural host." Virology (2002) 296:177-189. Deletion of such regions within a coronavirus such as SARS can be achieved, for example, by reverse genetics or "targeted recombination" (See, e.g., Masters, P. S., "Reverse genetics of the largest RNA viruses", Adv. Virus Res. (1999) 53:245-264.

Methods of purification of attenuated virus are known in the art and may include one or more of, for instance gradient centrifugation and chromatography. See Gregersen "Herstellung von Virussimpfstoffen aus Zellkulturen" Chapter 4.2 in Pharmazeutische Biotecnologie (eds. O. Kayser and RH Mueller) Wissenschaftliche Verlagsgesellschaft, Stuttgart, 2000.

C. Split SARS Vaccines

The invention includes a composition comprising a split SARS virus formulation and methods for the manufacture thereof. This composition can be used as a prophylactic or therapeutic SARS virus vaccine.

Methods of splitting enveloped viruses are known in the art. Methods of splitting enveloped viruses are disclosed, for example, in WO 02/28422, incorporated herein by reference in its entirety, and specifically including the splitting agents and methods described therein. Methods of splitting influenza viruses are disclosed, for example, in WO 02/067983, WO 02/074336, and WO 01/21151, each of which is incorporated herein by reference in its entirety.

The splitting of the virus is carried out by disrupting or fragmenting whole virus, infectious (wild-type or attenuated) or non-infectious (for example inactivated), with a disrupting

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concentration of a splitting agent. The disruption results in a full or partial solubilisation of the virus proteins, altering the integrity of the virus.

Preferably, the splitting agent is a non-ionic or an ionic surfactant. Accordingly, the split SARS virus formulations of the invention may also comprise at least one non-ionic surfactant or detergent. Examples of splitting agents useful in the invention include: bile acids and derivatives thereof, non-ionic surfactants, alkylglycosides or alkylthioglycosides and derivatives thereof, acyl sugars, sulphobetaines, betains, polyoxyethylenealkylethers, N,N-dialkyl-Glucamides, Hecameg, alkylphenoxypolyethoxyethanols, quaternary ammonium compounds, sarcosyl, CTAB (cetyl trimethyl ammonium bromide) or Cetavlon.

Preferably, the ionic surfactant is a cationic detergent. Cationic detergents suitable for use in the invention include detergents comprising a compound of the following formula:

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R₁, R₂ and R₃ are the same or different and each signifies alkyl or aryl, or

 R_1 and R_2 , together with the nitrogen atom to which these are attached form a 5- or 6-membered heterocyclic ring, and

R₃ signifies alkyl or aryl, or

R₁, R₂ and R₃ together with the nitrogen atom to which these are attached, signify a 5- or 6-membered heterocyclic ring, unsaturated at the nitrogen atom.

R₄ signfies alkyl or aryl, and

X signifies an anion.

Examples of such cationic detergents are cetyltrimethylammonium salts, such as ceyltrimethylammonium bromide (CTAB) and myristyltrimethylammonium salt.

Additional cationic detergents suitable for use in the invention include lipofectine, lipofectamine, and DOT-MA.

Non-ionic surfactants suitable for use in the invention include one or more selected from the group consisting of the octyl- or nonylphenoxy polyoxyethanols (for example the commercially available Triton series), polyoxyethylene sorbitan esters (Tween series) and polyoxyethylene ethers or esters of the general formula:

wherein n is 1-50, A is a bond or -C(O)-, R is C_{1-50} alkyl or phenyl $C_{1.50}$ alkyl; and combinations of two or more of these.

The invention comprises a method of preparing a split SARS virus comprising contacting the SARS virus with a sufficient amount of splitting agent to disrupt the viral envelope. The loss of integrity after splitting renders the virus non-infectious. Once the disrupted viral envelope proteins are generally no longer associated with whole intact virions, other viral proteins are preferably fully or partially solubilized and are therefore not associated, or only in part associated, with whole intact virions after splitting.

The method of preparing a split SARS virus may further comprise removal of the splitting agents and some or most of the viral lipid material. The process may also include a number of different filtration and/or other separation steps such as ultracentrifugation, ultrafiltration, zonal centrifugation and chromatographic steps in a variety of combinations. The process may also optionally include an inactivation step (as described above) which may be carried out before or after the splitting. The splitting process may be carried out as a batch, continuous, or semi-continuous process.

Split SARS virus vaccines of the invention may include structual proteins, membrane fragments and membrane envelope proteins. Preferably, the split SARS virus preparations of the invention comprise at least half of the viral structural proteins.

One example of a method of preparing a split SARS virus formulation includes the following steps:

- (i) propagation of the SARS virus in cell culture, such as MRC-5 cells (ATCC CCL-171), WI-38 cells (ATCC CCL-75), fetal rhesus kidney cells or vero cells (See the discussion in Section A, above, regarding culture of SARS virus);
 - (ii) harvesting of SARS virus-containing material from the cell culture;
 - (iii) clarification of the harvested material to remove non-SARS virus material;
 - (iv) concentration of the harvested SARS virus;
 - (v) separation of the whole SARS virus from non-virus material;
- (vi) splitting of the whole SARS virus using a suitable splitting agent in a density gradient centrifugation step; and
 - (vii) filtration to remove undesired materials.

The above steps are preferably performed sequentially.

The clarification step is preferably performed by centrifugation at a moderate speed. Alternatively, a filtration step may be used for example with a $0.2\mu m$ membrane.

The concentration step may preferably employ an adsorption method, for instance, using CaHPO4. Alternatively, filtration may be used, for example ultrafiltration.

A further separation step may also be used in the method of the invention. This further separation step is preferably a zonal centrifugation separation, and may optionally use a sucrose gradient. The sucrose gradient may further comprise a preservative to prevent microbial growth.

The splitting step may also be performed in a sucrose gradient, wherein the sucrose gradient contains the splitting agent.

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The method may further comprise a sterile filtration step, optionally at the end of the process. Preferably, there is an inactivation step prior to the final filtration step.

Methods of preparing split SARS virus formulations may further include treatment of the viral formulation with a DNA digesting enzyme. These enzymes may be used to reduce the level of host cell DNA in the viral purification process. DNA digesting enzymes for use in cell culture are known in the art and include, for example, Benzonase.

Treatment of the SARS virus formulation with a DNA digesting enzyme may occur at any time in the purification and splitting process. Preferably, however, the SARS virus formulation is treated with a DNA digesting enzyme prior to use of a detergent. Still more preferably, the SARS virus formulation is treated with a DNA digesting enzyme, such as Benzonas, prior to treatment with a cationic detergent, such as CTAB.

Methods of purification of split virus are known in the art. See JP Gregersen "Herstellung von Virussimpfstoffen aus Zellkulturen" Chapter 4.2 in Pharmazeutische Biotecnologie (eds. O. Kayser and RH Mueller) Wissenschaftliche Verlagsgesellschaft, Stuttgart, 2000.

The invention includes a split vaccine composition comprising a prophylactically effective amount of SARS viral antigen, preferably spike or an immunogenic fragment thereof. The SARS viral antigen is preferably present in a concentration amount of 0.1 to 50 μ g antigen/dose, more preferably 0.3 to 30 μ g antigen/dose. Still more preferably, the antigen is about 15 μ g/dose.

In one embodiment, a lower concentration of SARS viral antigen is used in split vaccine compositions of the invention. Such lower concentration vaccines may optionally comprise an adjuvant to boost the host immune response to the antigen. In such a "low dose" vaccine, the SARS viral antigen is preferably present in a concentration of less than 15 μ g antigen/dose, (i.e., less than 10, 7.5, 5 or 3 μ g antigen/dose.

D. Subunit SARS Vaccines

The invention includes a composition comprising an isolated or purified SARS viral antigen or a derivative thereof. The composition may further comprise one or more adjuvants.

SARS viral antigens can be isolated or purified from a SARS virus grown in cell culture.

Alternatively, SARS viral antigens can be recombinantly produced by methods known in the art.

The SARS viral antigens used in the invention can be produced in a variety of different expression systems which are known in the art; for example those used with mammalian cells, baculoviruses, bacteria, and yeast. Such expression systems will typically use polynucleotides encoding the viral antigens of the invention. Such sequences can be obtained using standard techniques of molecular biology, including translating the amino acid sequences listed herein. Accordingly, the invention includes polynucleotides encoding for the viral antigens of the

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invention. In addition, the viral antigens of the invention can be produced (at least in part, preferably in whole) via synthetic chemistry methods.

Insect cell expression systems, such as baculovirus systems, are known to those of skill in the art and described in, e.g., Summers and Smith, Texas Agricultural Experiment Station
Bulletin No. 1555 (1987). Materials and methods for baculovirus/insert cell expression systems are commercially available in kit form from, inter alia, Invitrogen, San Diego CA. Similarly, bacterial and mammalian cell expression systems are also known in the art and described in, e.g., Yeast Genetic Engineering (Barr et al., eds., 1989) Butterworths, London.

A number of appropriate host cells for use with the above systems are also known. For

example, mammalian cell lines are known in the art and include immortalized cell lines available from the American Type Culture Collection (ATCC), such as, but not limited to, Chinese hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (e.g., Hep G2), Madin-Darby bovine kidney ("MDBK") cells, as well as others. Mammalian sources of cells include, but are not limited to, human or non-human primate (e.g., MRC-5 (ATCC CCL-171), WI-38 (ATCC CCL-75), fetal rhesus lung cells (ATCC CL-160), human embryonic kidney cells (293 cells, typically transformed by sheared adenovirus type 5 DNA), VERO cells from monkey kidneys), horse, cow (e.g., MDBK cells), sheep, dog (e.g., MDCK cells from dog kidneys, ATCC CCL34 MDCK (NBL2) or MDCK 33016, deposit number DSM ACC 2219 as described in WO 97/37001), cat, and rodent (e.g., hamster cells such as BHK21-F, HKCC cells, or Chinese hamster ovary cells (CHO cells)), and may be obtained from a wide variety of developmental stages, including for example, adult, neonatal. fetal, and embryo.

Similarly, bacterial hosts such as E. coli, Bacillus subtilis, and Streptococcus spp., will find use with the present expression constructs. Yeast hosts useful in the present invention include, inter alia, Saccharomyces cerevisiae, Candida albicans, Candida maltosa, Hansenual polymorpha, Kluyveromyces fragilis, Kluyveromyces lactis, Pichia guillerimondii, Pichia pastoris, Schizosaccharomyces pombe and Yarrowia lipolytica. Insect cells for use with baculovirus expression vectors include, inter alia, Aedes aegypti, Autographa californica, Bombyx mori, Drosophila melanogaster, Spodoptera frugiperda, and Trichoplusia ni.

Nucleic acid molecules comprising nucleotide sequences of the viral antigens or antibodies of the invention can be stably integrated into a host cell genome or maintained on a stable episomal element in a suitable host cell using various gene delivery techniques well known in the art. See., e.g., US Patent No. 5,399,346.

Depending on the expression system and host selected, the molecules are produced by growing host cells transformed by an expression vector under conditions whereby the protein is expressed. The expressed protein is then isolated from the host cells and purified. If the expression system secretes the protein into growth media, the product can be purified directly

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from the media. If it is not secreted, it can be isolated from cell lysates. The selection of the appropriate growth conditions and recovery methods are within the skill of the art.

The invention includes a composition comprising an isolated or purified SARS viral antigen or a derivative thereof. The invention also includes a composition comprising at least two isolated or purified SARS viral antigens or derivatives thereof, which have been co-purified or purified separately and then combined. In one embodiment, the SARS viral antigen is a spike (S) protein. In yet another embodiment, the SARS viral antigen is a nucleocapsid (N) protein, a membrane (M) glycoprotein, or an envelope (E) protein. Preferably, the SARS viral antigen is present in the composition in a purity greater than 75% (e.g., 78%, 80%, 82%, 85%, 88%, 90%, 92%, 95%, 98%).

The invention includes a vaccine composition comprising a prophylactically effective amount of SARS viral antigen, preferably spike or an immunogenic fragment thereof. The SARS viral antigen is preferably present in a concentration amount of 0.1 to 50 μ g antigen/dose, more preferably 0.3 to 30 μ g antigen/dose. Still more preferably, the antigen is about 15 μ g/dose.

In one embodiment, a lower concentration of SARS viral antigen is used in vaccine compositions of the invention. Such lower concentration vaccines may optionally comprise an adjuvant to boost the host immune response to the antigen. In such a "low dose" vaccine, the SARS viral antigen is preferably present in a concentration of less than 15 μ g antigen/dose, (i.e., less than 10, 7.5, 5 or 3 μ g antigen/dose.

The following example illustrates a method of preparing a SARS virus spike (S) protein subunit vaccine.

SARS virus S antigen may be isolated and purified from a variety of sources and using a variety of methods, including, but not limited to, S antigen expressed in cultured eukaryotic cells (e.g., mammalian cells, such as VERO, CHO) or bacteria (e.g., E. coli). Expression of may be achieved by a variety of means, such as, for example, from SARS virus infected cell culture or cell culture supernatants, from cultured cells stably transformed with a DNA expression cassette encoding the SARS virus S protein (e.g., RNA polymerase II promoter operably linked to a SARS virus S gene), or from cultured cells infected with a replication-competent or replication-incompetent virus-based expression vector (e.g., adenovirus vector, poxvirus vector, alphavirus vector, retrovirus vector) encoding the SARS virus S protein, as a means to eliminate the need to work with infectious SARS virus.

1. Subunit SARS Vaccines Produced from SARS Virus Cultures

The SARS virus may be grown in cultured mammalian celle, such as VERO cells, then separated from the cultured cells. A SARS viral antigen, such as the S protein, can then be solubilized and separated from the SARS virus, and further isolated and purified.

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In one example, the SARS virus may be produced as described in the Inactivated SARS vaccine examples, then the desired SARS antigen, such as spike protein, may be further purified from the end product using techniques known in the art.

In another example, a SARS subunit vaccine may be produced as follows. SARS virus may be produced using a desired mammalian cell line on microcarrier beads in large, controlled fermentors. For example, vaccine quality African Green Monkey kidney cells (VERO cells) at a concentration of 10⁵ cells/mL are added to 60 to 75 L of CMRL 1969 media, pH 7.2, in a 150 L bioreactor containing 360 g of Cytodex-1 microcarrier beads and stirred for 2 hours. Additional CMRL 1969 is added to give a total volume of 150 L. Fetal bovine serum (FBS) is added to a final concentration of 3.5%. Glucose is added to a final concentration of 3.0 g/L and glutamine is added to a final concentration of 0.6 g/L. Dissolved oxygen, pH, agitation and temperature are controlled, and cell growth, glucose, lactate and glutamine levels are monitored. When cells are in logarithmic phases usually on days 3 to 4 reached a density of about 1.0-2.5x106 cells/mL, the culture medium is drained from the fermentor and 120 L of CMRL 1969, pH 7.2 (no FBS) is added and the culture stirred for 10 minutes. The draining and filling of the fermentor is usually repeated once but could be repeated up to three times. After washing the cells, the fermentor is drained and 50 L of CMRL 1969 containing 0.1% (v/v) FBS is added. The SARS virus inoculum is added at a multiplicity of infection (m.o.i.) of 0.001 to 0.01. Trypsin may be added to promote efficient infection. Additional CMRL 1969 with 0.1% FBS is added to give a final volume of 150 L. Incubation is continued at 34 C. One viral harvest is obtained from a single fermentor lot, typically at 2-7 days post-infection. Multiple harvests from a single fermentation may also be obtained.

The isolation and purification of S protein may be effected by a variety of means, as described below. For example, collecting S protein-containing flow-through from ion exchange chromatography of solubilized SARS virus envelope proteins; loading the flow through onto a hydroxyapatite matrix, and selectively eluting the S protein from the hydroxyapatite matrix. The selectively eluted S protein may be further concentrated by tangential flow ultrafiltration.

Alternatively, the isolation and purification may be effected by collecting S protein-containing flow-through from ion exchange chromatography of the solubilized SARS virus envelope proteins; loading the flow through onto a hydroxyapatite matrix and collecting an S protein-containing flow through, selectively removing detergent used in the solubilization step from the hydroxyapatite matrix flow through to provide isolated and purified S protein. The isolated and purified S protein may be subsequently concentrated by tangential flow ultrafiltration

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Nucleic acid contaminants may be removed from the isolated and purified S protein by treatment with a nucleic acid degrading agent as described above in the Inactivation section. Preferably, the nucleic acid degrading agent is a nuclease, such as for example. Benzonase.

The isolated and purified S protein may be applied to a gel filtration medium and the S protein subsequently collected therefrom to separate the S protein from contaminants of other molecular weights.

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Alternatively, the isolation and purification may be effected by loading S protein on a first ion-exchange medium while permitting contaminants to pass through the medium, eluting the S protein from the first ion-exchange medium, to separate the S protein from contaminants of other molecular weights. The eluted S protein is applied to a second ion-exchange medium while allowing contaminants to pass through the second ion-exchange medium. The S protein is subsequently eluted therefrom, to provide the isolated and purified S protein. The eluted S protein may be concentrated by tangential flow ultrafiltration.

Alternatively, substantially pure SARS virus S protein suitable for use as an immunogen in a subunit vaccine formulation may be prepared from infected cell lysates, such as for example using a non-denaturing detergent buffer containing 1% Triton X-100 and deoxycholate to lyse infected cells. The cell lysates are clarified by centrifugation and S protein is purified from the cell lysates by immunoaffinity purification. A monoclonal antibody against the S protein is generated and coupled to beads and a column is constructed with those beads. SARS-infected cell lysates are applied to the column, and the column is washed with PBS containing 0.1% Triton X-100. Protein bound to the column is eluted with 0.1M glycine, pH 2.5, 0.1% Triton X-100. Elution samples are buffered, such as for example, with Tris, and analyzed for the presence of protein. Fractions containing the protein are pooled and dialyzed against PBS

As discussed above, the present invention includes isolated and purified S protein of SARS virus. In one example, the virus is grown on a vaccine quality cell line, such as VERO cells, and the grown virus is harvested. The virus harvest is filtered and then concentrated typically using tangential flow ultrafiltration using a membrane of desired molecular weight cut-off and diafiltered. The virus harvest concentrate may be centrifuged and the supernatant discarded. The pellet from the centrifugation then is detergent extracted to solubilize the S protein, for example, by resuspending the pellet to the original harvest concentrate volume in an extraction buffer containing a detergent such as a non-ionic detergent including TRITON X-100.

Following centrifugation to remove non-soluble proteins, the S protein extract is purified by chromatographic procedures. The extract may first be applied to an ion exchange chromatography column such as a TMAE-fractogel or S-fractogel column equilibrated to permit the S protein to flow through while impurities are retained on the column.

Next, the flow through may be loaded onto a hydroxyapatite column, equilibrated to permit binding of the S protein to the matrix and to permit contaminants to pass from the column. The bound S protein is then eluted from the column by a suitable elutant. The resulting purified solution of S protein may be further processed to increase its purity. The eluate first may be concentrated by tangential flow ultrafiltration using a membrane of desired molecular weight cut-off. The filtrate may be contacted with a polyethylene glycol of desired molecular weight, for example, about 6000 to 8000, to precipitate the protein. Following centrifugation and discard of the supernatant, the pellet may be resuspended in PBS and dialyzed to remove the polyethylene glycol. Finally, the dialyzed solution of S protein may be sterile filtered. The sterile filtered solution may be adsorbed onto alum. The polyethylene glycol precipitation and resuspension purification step may be effected at an earlier stage of the purification operation, if desired.

Alternatively, SARS virus is recovered following growth and harvesting of the virus, and a concentrate obtained such as, for example using PEG precipitation or tangential flow filtration. The virus is contacted with detergent to solubilize the S proteins. Following centrifugation, the supernatant is recovered to further purification of the S protein and the non-soluble proteins discarded.

The supernatant is applied to an ion exchange chromatography column, such as a TMAE-fractogel or S-fractogel column, suitably equilibrated to permit retention of the S protein on the column. The S protein is eluted from the ion-exchange column under suitable conditions. The eluate then may be passed through a gel filtration column, such as a Sephacryl S-300 column, to separate the S protein from contaminants of other molecular weights. A hydroxyapatite column may be employed in place of the Sephacryl column.

The S protein may be eluted from the column to provide a purified solution of S protein. The eluate may be concentrated by tangential flow ultrafiltration using a membrane of desired molecular weight cut-off. The concentrated S protein solution then may be sterile filtered.

Alternatively, viral harvests may be concentrated by ultrafiltration and the concentrated viral harvests may be subjected to an initial purification step, for example, by gel filtration chromatography, polyethylene glycol precipitation or Cellufine sulfate chromatography. The purified virus may then be detergent extracted to solubilize the S protein. Following solubilization of the S protein, the supernatant may be loaded onto an ion-exchange column such as Cellufine sulfate chromatography column equilibrated to permit the protein to bind to the column while permitting contaminants to flow through. Similarly, a TMAE-fractogel or S-fractogel column may be used in place of the Cellufine sulfate column. The two columns also may be combined in sequential purification steps. The S protein is eluted from the columns to

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provide a purified solution of the protein. This solution may be concentrated by tangential flow ultrafiltration using a membrane of desired molecular weight cut-off and diafiltered.

Specifically, in one method of S protein purification, the virus harvest concentrate is centrifuged at 28,000 x g for 30 minutes at 4 C. The supernatant is discarded and the pellet resuspended in extraction buffer consisting of 10 mM Tris-HCl, pH 7.0, 150 mM NaCl, 2% (w/v) Triton X-100 to the original harvest concentrate volume. Pefabloc is added to a final concentration of 5 mM. The suspension is stirred at room temperature for 30 minutes. The supernatant, containing the soluble S protein, is clarified by centrifugation at 28,000 x g for 30 minutes at 4 C. A TMAE--Fractogel column is equilibrated with 10 mM Tris-HCl, pH 7.0, 150 mM NaCl containing 0.02% Triton X-100. The Triton X-100 supernatant, containing the soluble S protein, is loaded directly onto the TRAE-Fractogel column. The total volume added plus 2 bed volumes of 10 mM Tris-HCl, pH 7.0, 150 mM NaCl containing 0.02% Triton X-100 are collected. The TMAE--Fractogel flow-through containing S protein is diluted 3-fold with 10 mM Tris-HCl, pH 7.0, containing 0.02% Triton X-100.

An hydroxyapatite column is equilibrated with 10 mM Tris-HCl, pH 7.0, 50 mM NaCl, 0.02% Triton X-100. After loading the TMAE flow-through, the column is washed with 2 column volumes of 10 mM Tris-HCl, pH 7.0, 50 mM NaCl, 0.02% Triton X-100 followed by 4 column volumes of 5 mM sodium phosphate, pH 7.0, 1M NaCl, 0.02% Triton X-100. The proteins are eluted with 4 column volumes of 20 mM sodium phosphate, pH 7.0, 1M NaCl, 0.02% Triton X-100. Fractions are collected based on A280 and the protein content and antigen concentrations are measured. The purified S protein is ultrafiltered by tangential flow ultrafiltration using a 300 kDa NMWL membrane.

2. Recombinant Production of Subunit SARS Vaccines

As discussed above, SARS virus proteins may be produced by recombinant expression.

Host cells suitable for recombinant expression include bacterial, mammalian, insect, yeast, etc.

Recombinant expression may be used to produce a full length SARS protein, a fragment thereof, or a fusion therewith.

Fusion peptides may be used to facilitate the expression and purification of the recombinant SARS protein. For example, recombinant production of the SARS polypeptides can be facilitated by the addition a tag protein to the SARS antigen to be expressed as a fusion protein comprising the tag protein and the SARS antigen. Such tag proteins can facilitate purification, detection and stability of the expressed protein. Tag proteins suitable for use in the invention include a polyarginine tag (Arg-tag), polyhistidine tag (His-tag), FLAG-tag, Strep-tag, c-myc-tag, S-tag, calmodulin-binding peptide, cellulose-binding domain, SBP-tag,, chitin-binding domain, glutathione S-transferase-tag (GST), maltose-binding protein, transcription termination anti-termination factor (NusA), E. coli thioredoxin (TrxA) and protein disulfide

isomerase I (DsbA). Preferred tag proteins include His-tag and GST. A full discussion on the use of tag proteins can be found at Terpe et al., "Overview of tag protein fusions: from molecular and biochemical fundamentals to commercial systems", Appl Microbiol Biotechnol (2003) 60:523-533.

After purification, the tag proteins may optionally be removed from the expressed fusion protein, *i.e.*, by specifically tailored enzymatic treatments known in the art. Commonly used proteases include enterokinase, tobacco etch virus (TEV), thrombin, and factor X...

Accordingly, the invention further includes a SARS virus subunit vaccine comprising a fusion protein. Preferably, the fusion protein comprises a first amino acid sequence encoded by a SARS virus polynucleotide sequence. SARS virus polynucleotide sequences which may encode said first amino acid sequence include one or more of the SARS virus polynucleotide sequences identified in this application and fragments thereof.

The fusion protein may comprise an amino acid sequence of a SARS virus protein or a fragment thereof. Said SARS virus protein may be selected from one or more of the group consisting of the following SARS virus proteins: P28, P65, Nsp1, Nsp2 (3CL protease), Nsp3, Nsp3, Nsp4, Nsp 5, Nsp6, Nsp 7, Nsp 8, Nsp 9 (RNA polymerase), Nsp 10 (helicase), Nsp 11, Nsp 12, Nsp 13, Spike, Orf 3, Orf 4, Envelope, Matrix, Orf 7, Orf 8, Orf 9, Orf 10, Orf 11, Nucleocapsid and Orf 13.

In one embodiment, the fusion protein comprises a first amino acid sequence comprising a SARS virus antigen or a fragment thereof. Said SARS virus amino acid sequence may comprise one or more of the T-epitope sequences identified above.

Preferably, the fusion protein comprises an amino acid sequence of a SARS virus spike protein, or a fragment thereof. Specific fragments of the spike protein which may be used in the fusion protein include the S1 domain and the S2 domain. Further fragments of the spike protein which may be used in the fusion protein include regions of each of the S1 and S2 domains, including the receptor binding region of the S1 domain, the oligomerization domain regions of the S2 domain, the leucine zipper regions of the S2 domain, the membrane anchor region of the S2 domain, the hydrophobic domain region of the S2 domain, the cystein-rich domain region of the S2 domain, and the cytoplasmic tail region of the S2 domain. (See FIGURE 19). Amino acid sequences of the Spike protein corresponding to these regions can be identified by those skilled in the art, including, for example, using the functional predictions set forth earlier in the application (predicted transmembrane helices, predicted N-terminus signaling regions, predicted coiled-coil regions, etc.) as well as by homology comparison to the sequences of other known Coronaviruses (See FIGURES 4F and 5).

The fusion protein may further comprise a second amino acid sequence. Said second amino acid sequence may comprise a polypeptide sequence which facilitates protein expression

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or purification, preferably one of the tag sequences discussed above. Alternatively, said second amino acid sequence may comprise a second amino acid sequence from a SARS virus. Alternatively, said second amino acid sequence may comprises an amino acid sequence from another virus or bacteria, including one or more of the viruses or bacteria identified in Section I, below.

Said second amino acid sequence may comprise an amino acid sequence from another respiratory virus. Said second amino acid sequence may comprise an amino acid sequence from a virus selected from the group consisting of coronavirus, influenza virus, rhinovirus, parainfluenza virus (PIV), respiratory syncytial virus (RSV), adenovirus, and metapneumovirus.

In one embodiment, said second amino acid sequence may comprise an amino acid sequence from an adjuvant, including one or more of the adjuvants identified in section I. below.

In one embodiment, the invention includes a fusion protein comprising an amino acid sequence of a SARS virus spike protein or a fragment thereof. The fusion protein may further comprise a second amino acid sequence comprising an amino acid sequence selected from the group consisting of a second SARS virus protein, a non-SARS virus protein, a bacterial protein, and an adjuvant.

(a) Bacterial Expression of Subunit SARS Vaccines

In one embodiment, bacterial host cells are used for recombinant expression of SARS virus proteins. Bacterial host cells suitable for use in the invention include, for example, E. coli, Bacillus subtilis, and Streptococcus spp.

The SARS viral protein may be modified to facilitate bacterial recombinant expression. In particular, the SARS spike protein may be modified to facilitate transport of the spike protein to the surface of the bacterial host cell.

Applicants have discovered that there is strong structural homology between the SARS virus spike protein and the NadA protein of *Neisseria meningitidis*. Both proteins have an N-terminal globular "head" domain (amino acids 24-87), an intermediate alpha-helix region with high propensity to form coiled-coil structures (amino acids 88-350), and a C-terminal membrane anchor domain formed by four amphipatix transmembrane beta strands (amino acids 351-405 of NadA). In addition, a leucine zipper motive is present within the coiled-coil segment. See, FIGURE 19 depicting the SARS spike protein structure Comanducci *et al.*, "NadA, a Novel Vaccine Candidate of Neisseria meningitidis", J. Exp. Med. 195 (11): 1445-1454 (2002). In addition, a leucine zipper motif of NadA is present within the coiled-coil segment. The NadA protein also forms high molecular weight surface-exposed oligomers (corresponding to three or four monomers) anchored to meningococcal outer membrane.

When the NadA protein is expressed in E. coli, the full-length protein is assembled in oligomers anchored to the outer membrane of E. coli, similar to the way the protein is presented

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in meningococcus. The NadA protein devoid of the predicted membrane anchor domain is then secreted into the culture supernatant. This secreted protein is soluble and still organized in trimers.

The invention therefore includes a fusion protein comprising an amino acid sequence of a SARS virus spike protein or a fragment thereof and a second amino acid sequence of a bacterial adhesion protein or a fragment thereof. Preferably, said adhesion protein is selected from the group consisting of NadA, YadA (of enteropathogenic Yersinia), and UspA2 (of Moraxella catarrhalis). Additional NadA-like proteins include serum resistance protein DsrA of Haemophilus ducreyi, the immunoglobulin binding proteins EibA, C, D, and F of E. coli, outer membrane protein 100 of Actinobacillus actinomycetemcomitans, the saa gene carried on the large virulence plasmid present in shiga toxigenic strains of E. coli (STEC), and each of the bacterial adhesion proteins described in U.K. Patent Application No. 0315022.4, filed on June 26, 2003, each of which are specifically incorporated herein by reference.

Preferably, said adhesion protein comprises NadA or a fragment thereof.

Such fusion proteins may be used to facilitate recombinant expression of immunogenic portions of SARS surface antigens, such as spike. These fusion constructs may also allow the SARS S1 and/or S2 domains to adapt to a native confirmation. These fusion proteins are also able to oligomerize and form dimers or trimers, allowing the S1 and/or S2 domains to associate and adapt conformations as in the native SARS spike protein. Further, these expression constructs facilitate surface exposure of the SARS spike protein.

The fusion proteins of the invention preferably comprise a leader peptide from a NadA like protein, preferably NadA, a polypeptide from the immunogenic "head" region of the spike protein, and a stalk region from either the NadA like protein or the Spike protein. During expression and processing of the fusion protein, one or more amino acids may be cleaved off or removed, such as, i.e., the leader peptide or a membrane anchor domain.

The stalk regions facilitate oligomerization of the expression protein. Optionally, the fusion proteins of the invention further include an anchor region of a NadA like protein. This anchor region allows the expression fusion protein to anchor and assemble on the bacterial cell surface.

The fusion proteins of the invention include the following constructs:

(i) the NadA leader peptide (optionally also including the first 6 amino acids of the mature NadA protein to facilitate processing of the leader peptide and appropriate maturation of the protein) followed by the Spike S1 domain. Preferably, this construct comprises amino acids 1-29 of NadA (corresponding to the NadA leader peptide and the first 6 amino acids of the mature NadA protein, as shown in FIGURE 22 and as set forth below) followed by amino acids 14-662

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of a SARS virus Spike protein (corresponding to the S1 domain, see FIGURE 19 and SEQ ID NO: 6042 and as set forth below). Specifically, construct (i) comprises SEO ID NO: 7302.

- (ii) the NadA leader peptide (optionally also including the first 6 amino acids of the mature NadA protein to facilitate processing of the leader peptide and appropriate maturation of the protein) followed by the Spike S1 domain, followed by the stalk and anchor membrane domains of NadA. Preferably, this construct comprises amino acids 1-29 of NadA (corresponding to the NadA leader peptide and the first 6 amino acids of the mature NadA protein, as shown in FIGURE 22 and as set forth below) followed by amino acids 14-662 of a SARS virus Spike protein (corresponding to the S1 domain, see FIGURE 19 and SEQ ID NO: 6042 and as set forth below) followed by amino acids 88-405 of NadA (corresponding to the stalk and the anchor membrane domains). Specifically, construct (ii) comprises SEQ ID NO: 7303.
- (iii) the NadA leader peptide (optionally also including the first 6 amino acids of the mature NadA protein) followed by a SARS virus Spike S1 domain, followed by the NadA stalk domain. Preferably, this construct comprises amino acids 1-29 of NadA followed by amino acids 14-662 of a SARS virus Spike protein (corresponding to the S1 domain), followed by amino acids 88-350 of NadA (corresponding to the stalk domain). Specifically, construct (iii) comprises SEQ ID NO: 7304.
- (iv) the NadA leader peptide (optionally also including the first 6 amino acids of the mature NadA protein), followed by a SARS virus Spike S1 and S2 domain (excluding the putative transmembrane region), followed by the anchor domain of NadA. Preferably, this construct comprises amino acids 1-29 of NadA, followed by amino acids 14-1195 of a SARS virus Spike protein (corresponding to S1 and S2, excluding the putative transmembrane region), followed by amino acids 351-405 of NadA (corresponding to the NadA anchor domain). Specifically, construct (iv) comprises SEQ ID NO: 7305. Alternatively, the NadA anchor domain may comprise amino acids 332 405 of NadA.
- (v) the NadA leader peptide (optionally also including the first 6 amino acids of the mature NadA protein), followed by a SARS virus Spike S1 and S2 domain (exclusing the putative transmembrane region). Preferably, this construct comprises amino acids 1-29 of NadA, followed by amino acids 14-1195 of a SARS virus Spike protein. Specifically, construct (v) comprises SEQ ID NO: 7306.

In each of constructs (i) to (v), the first 23 amino acids are the NadA leader peptide, and the GS dipeptide at residues 679-680 arises from the insertion of a restriction enzyme site.

In constructs (i), (ii) and (iii), the NadA "head" is replaced by the Spike S1 domain, and the fusion proteins are anchored to the outer membrane of *E.coli* or secreted in the culture supernatant, respectively. In constructs (iv) and (v), the "head" and "stalk" domains of NadA are

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replaced by S1 and S2 Spike domains; also in this case, the two fusion proteins are anchored to the outer membrane of *E. coli* or secreted in the culture supernatant, respectively.

Accordingly, the invention further includes a fusion protein comprising an amino acid sequence of a SARS virus spike protein or a fragment thereof and a second amino acid sequence of a bacterial adhesion protein or a fragment thereof. Preferably, amino acids corresponding to the "head" of the adhesion protein are replaced by amino acids corresponding to a SARS virus Spike S1 domain. Alternatively, the amino acids corresponding to the "head" and "stalk" domains of the bacterial adhesion protein are replaced by amino acids corresponding to the SARS virus spike protein S1 and S2 domains.

As discussed above and shown in Figure 19, the S1 domain of the Spike protein is identified as the globular receptor binding "head" region. The S1 domain of the Spike protein preferably comprises about amino acids 14-662 of SEQ ID NO: 6042. The S1 domain may comprise a shorter amino acid sequence, wherein amino acids are removed from either the N-terminal or C-terminal regions. Preferably, 3, 5, 7, 9, 13, 15, 20 or 25 amino acids are removed from either the N-terminal or C-terminal regions. The S1 domain further includes amino acid sequences having sequence identity to the S1 region of SEQ ID NO: 6042. An example of the S1 domain is SEO ID NO: 7307:

As discussed above and shown in Figure 19, the S2 domain of the Spike protein is identified as the "stalk" region. The "stalk" region comprises oligomerization domain regions, a leucine zipper domain regions, membrane anchor regions, hydrophobic domain regions, cystein-rich domain region and a cytoplasmic tail region. The S2 domain of the Spike protein preferably excludes the transmembrane region and comprises about amino acids 663-1195 of SEQ ID NO: 6042. The S2 domain may comprise a shorter amino acid sequence, wherein amino acids are removed from either the N-terminal or C-terminal regions. Preferably, 3, 5, 7, 9, 13, 15, 20 or 25 amino acids are removed from either the N-terminal or C-terminal regions. The S2 domain further includes amino acid sequences having sequence identity to the S2 region of SEQ ID NO: 6042. An example of the S1 domain (with the transmembrane region excluded) is SEQ ID NO: 7308.

An example of the NadA protein described above is SEQ ID NO: 7309. As discussed above, the leader sequence of NadA used in the fusion protein preferably comprises about the first 29 amino acids of NadA (including a leader sequence with about 6 amino acids of the NadA head protein). Examples of such a leader sequences are set forth as SEQ ID NOS: 7310 and 7311 below. The fusion protein may use a leader sequence comprising a shorter amino acid sequence, wherein amino acids are removed from either the N-terminal or C-terminal end of the sequence. The leader sequence used in the fusion protein may also include an amino acid

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sequences having sequence identity to SEQ ID NO: 7310 or SEQ ID NO: 7311. Preferably, the leader sequence comprises SEQ ID NO: 7311.

Optionally, the fusion peptide comprises about the first 6 amino acids of the mature NadA protein to facilitate processing of the leader peptide and appropriate maturation of the protein.

An examples of the first 6 amino acids of a mature NadA proteins is SEO ID NO: 7312...

As discussed above, the stalk and anchor sequences of NadA used in the fusion protein preferably comprise about amino acids 88-405 of NadA. An example of an amino acid sequence comprising NadA stalk and anchor regions is set forth below as SEQ ID NO: 7313 below. An example of an amino acid sequence comprising a NadA stalk region (without the anchor region) is set forth as SEQ ID NO: 7314 below. An example of an amino acid sequence comprising a NadA anchor region is set forth as SEQ ID NO: 7315 below. The fusion protein may use a stalk (and/or anchor) sequence comprising a shorter amino acid sequence, wherein amino acids are removed from either the N-terminal or C-terminal regions. Preferably, 1, 2, 3, 4, 5, 6, 7, 8 or 9 amino acids are removed from either the N-terminal or C-terminal end of the sequence. The leader sequence used in the fusion protein may also include an amino acid sequences having sequence identity to the SEQ ID NO: 7313.

The fusion proteins of the invention, including those described above, may be prepared, for example, as follows. Single fragments (such as the regions described above) may be amplified by PCR using the oligonucleotide primers set forth in the Table below. (S1_L refers to the Spike protein fused to the leader peptide of NadA; S2 refers to the stalk region of the Spike protein, with and without the stop codon). The oligonucleotides were designed on the basis of the DNA sequence of NadA from N. meningitidis B 2996 strain and of Spike from SARS virus isolate FRA1. Each oligonucleotide includes a restriction site as a tail in order to direct the cloning into the expression vector pET21b.

		SEQ ID NO:	Restriction site
$S1_L$	For	7316	NdeI
S1 _L	Rev	7317	BamHI
S2	For	7318	BamHI
S2	Rev	7319	HindⅢ
S2-stop	Rev	7320	XhoI
NadA ₈₈	For	7321	BamHI
NadA ₃₅₀	Rev	7322	XhoI
NadA ₃₃₂	For	7323	HindIII
NadA ₄₀₅	Rev	7324	XhoI

The single fragments are sequentially cloned into pET21b vector, in order to express the proteins under the control of inducible T7 promoter. The S1 domain of the Spike protein fused to the leader peptide of NadA (S1_L) was obtained by PCR using the primers S1_L-For and S1_L-Rev. The forward oligonucleotide primer contains the NdeI restriction sequence and the

sequence coding for the leader peptide of NadA plus the first 6 aminoacids of the mature protein. The PCR fragment was cloned as a NdeJ/BamHI fragment in the pET21b vector opened with the same restriction enzymes. This clone (pET-S1_L) was then used to sequentially clone the other different domains, as BamHI/XhoI, BamHI/HindIII or HindIII/XhoI fragments. BamHI and HindIII restriction sites introduce the aminoacids GS and KL, respectively.

The PCR amplification protocol was as follows: 200ng of genomic DNA from Neisseria meningitidis 2996 or 10 ng of plasmid DNA preparation (plasmid pCMVnew, containing the entire gene coding of the Spike protein), were used as template in the presence of 40μM of each oligonucletide primer, 400-800 μM dNTPs solution, 1x PCR buffer (including 1.5mM MgCl₂), 2.5 units TaqI DNA polymerase (using Perkin-Elmer AmpliTaQ or Invitrogen Platinum Pfx DNA polymerase).

After a preliminary 3 minute incubation of the whole mix at 95°C, each sample underwent a two-step amplification: the first 5 cycles were performed using the hybridisation temperature that excluded the restriction enzyme tail of the primer (Tm1). This was followed by 30 cycles according to the hybridisation temperature calculated for the whole length oligos (Tm2). Elongation times, performed at 68°C or 72°C, varied according to the length of the fragment to be amplified. The cycles were completed with a 10 minute extension step at 68°C or 72°C.

The amplified DNA was either loaded directly on agarose gel and the DNA fragment corresponding to the band of correct size was purified from the gel using the QiagenTM Gel Extraction Kit, following the manufacturer's protocol.

The purified DNA corresponding to the amplified fragment and the plasmid vectors' were digested with the appropriate restriction enzymes, purified using the QIAquick TM PCR purification kit (following the manufacturer's instructions) and ligation reactions were performed.

The ligation products were transformed into competent *E. coli* DH5a and screening for recombinant clones was performed by growing randomly-selected colonies and extracting the plasmid DNA using the Qiagen QIAprep Spin Miniprep Kit, following the manufacturer's instructions.

Recombinant plasmids were introduced into *E. coli* BL21(DE3) used as expression host. Single recombinant colonies were inoculated into LB + ampicillin and incubated at 37°C for 14-16 h. Bacteria were directly recovered by centrifugation (uninduced conditions) or diluted in fresh medium and grown at 37°C until OD $_{600}$ between 0.4-0.8. Protein expression was induced by addition of 1 mM Isopropyl-1-thio- β -D-galactopyranoside (IPTG) for three hours (induced conditions).

Whole cell lysates were obtained resuspending bacteria in SDS-sample buffer 1X and boiling for 5-10 min. Equal amounts of proteins were separated using NuPAGE (Invitrogen) or -131-

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BIORAD Gel System, according to the manufacturer's instructions. Proteins were revealed by Coomassie-blue staining or transferred onto nitrocellulose membranes for western blot analysis. Western blot was performed using a rabbit polyclonal anti-serum against purified NadA_{A351-405} (diluted 1:3000) and a secondary peroxidase-conjugate antibody (DAKO).

Results of the expression in E.coli of $S1_L$, $S1_L$ -NadA and $S1_L$ -NadA_{Aanchor} are shown in FIGURES 38 and 39. Schematics of the fusion constructs are shown in FIGURE 37.

Bacterial expression of the SARS viral antigens may also be used to prepare compositions comprising outer membrane vesicles wherein said outer membrane vesicles comprise one or more SARS viral antigens.

Outer Membrane Vesicles ("OMV"), also referred to as blebs, refer to vesicles formed or derived from fragments of the outer membrane of a Gram negative bacterium. OMVs typically comprise outer membrane proteins (OMPs), lipids, phospholipids, periplasmic material and lipopolysaccharide (LPS). Gram negative bacteria often shed OMVs during virulent infections in a process known as blebbing. OMVs can also be obtained from Gram negative bacteria via a number of chemical denaturation processes, such as detergent extraction. Synthetic OMVs or liposomes, comprising a lipid bilayer and typically enclosing an aqueous core, can also be prepared with the SARS viral antigens of the invention.

The OMVs of the invention are preferably lipid vesicles comprising a lipid bilayer surrounding an aquous core. Typically the lipid vesicles are of unilamellar structure (i.e., a single lipid bilayer surrounds the aquous core), although multilammelar lipid vesicles may also be used in the compositions of the invention. OMVs typically have sizes in the nanomolar to micromolar range, e.g., from 1 nM to 100 μ M, more typically from 10nM to 10 μ M and preferably from 30 nM to 1 μ M.

The OMVs of the invention are preferably prepared from gram negative bacteria. Gram negative bacteria are those bacteria that fail to resist decolorization in the commonly known Gram staining method. Gram negative bacteria are characterized by a complex multilater cell wall and often possess an outer layer polysaccharide capsule. Gram negative bacteria suitable for producing OMVs include, for example, species from Neisseria, Moraxella, Kingella, Acinetobacter, Brucella, Bordetella, Chlamydia, Porphyromonas, Actinobacillus, Borelia, Serratia, Campylobacter, Helicobacter, Haemophilus, Escherichia, Legionella, Salmonella, Pseudomonas and Yersinia.

The OMVs of the invention preferably comprise one or more SARS viral antigens or a fragment thereof. The SARS viral antigens may be recombinantly expressed in a Gram negative bacterial host cell and then harvested with the OMV.

Antigenic components, such as recombinantly expressed SARS viral antigens, may be located in any or all of the three main compartments of the lipid vesicles, including attached to

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either the interior or exterior surface of the lipid vesicle, for example via a membrane anchor domain, or attachment to a lipid moiety; inserted into the lipid bilayer, for example where the antigenic component is itself a hydrophobic or lipid based entity; or located within the aqueous center or core of the lipid vesicle.

Synthetically prepared OMVs, or liposomes, may be used in the invention. Such liposomes may comprise a number of different lipids and fatty acids. Suitable lipids for inclusion in liposomes of the invention include but are not limited to phophatidylinositol-(4,5)-diphosphate, phosphatidylserine, phosphatidylcholine, phosphatidylethanolamine, phosphatidyglycerol, cholesterol, beta-oleolyl-gamma-palmitoyl, lipopolysaccharides and galactocerbrosides.

Suitable means for extraction of OMVs from bacterial sources include deoxycholate extraction, Tris/HCl/EDTA extraction, and lithium acetate extraction. Preferably, the extraction process comprises a physical and/or chemical means to disrupt the bacterial cell outer membrane in order to release sufficient OMVs for purification and isolation. See, e.g., WO 03/051379.

The OMVs of the invention may be enriched and/or supplemented with antigenic components, such as SARS viral antigens, by methods known in the art, including, for example, direct combination in vitro where an energetic combination step can optionally be applied to facilitate integration of the antigenic component into a compartment of the liposome. Methods of energetic combination suitable for use in the invention include homogenization, ultrasonication, extrusion, and combinations thereof.

Preferably, the antigenic component, such as the SARS viral antigen, is recombinantly produced by the host cell from which the OMV is derived. In one embodiment, such OMVs are prepared by introducing nucleic acid sequence encoding for the SARS viral antigen into the recombinant host cell. Preferably the nucleic acid sequence encoding for the SARS viral antigen is controlled by a strong promoter sequence. Preferably, the nucleic acid sequence encoding the SARS viral antigen further comprises an outer-membrane targeting signal. For example, the nucleic acid sequence encoding the SARS viral antigen may be fused to a sequence encoding for a naturally occurring outer membrane protein of the bacterial host. Preferably, the nucleic acid sequence encoding the SARS viral antigen is fused to the signal peptide sequence of the naturally occurring outer membrane protein of the bacterial host.

Methods of preparing an optimizing OMVs for use in vaccines are disclosed in, for example Filip et al., J. Bact. (1973) 115: 717-722; Davies et al., J. Immunol. Method (1990) 143:215-225; and WO 01/09350.

In one embodiment, a bacterial host cell, such as *E. coli*, are transformed to express the SARS spike protein. As discussed above, the spike protein may be modified to facilitate bacterial expression and transport of the spike protein to the surface of the host cell. Each of the

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Spike/NadA fusion constructs discussed above may be used in the OMV preparations of the invention. Preferably, constructs comprising the spike S1 globular head domain fused to the stalk region of NadA are used to generate OMVs. The construct may optionally include the NadA leader peptide as well as the NadA anchor peptide. Schematic diagrams of these preferred OMV constructs are depicted in FIGURE 49.

Example 6 describes one method of preparing the OMVs of the invention.

(b) Mammalian Expression of Subunit SARS Vaccine

As discussed above, mammalian host cells may be used for recombinant expression of SARS virus proteins. Mammalian host cells suitable for use in the invention include, for example, Chinese hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (e.g., Hep G2), Madin-Darby bovine kidney ("MDBK") cells, as well as others. Mammalian sources of cells include, but are not limited to, human or non-human primate (e.g., MRC-5 (ATCC CCL-171), WI-38 (ATCC CCL-75), human embryonic kidney cells (293 cells, typically transformed by sheared adenovirus type 5 DNA), VERO cells from monkey kidneys (including, for example COS7 cells), horse, cow (e.g., MDBK cells), sheep, dog (e.g., MDCK cells from dog kidneys, ATCC CCL34 MDCK (NBL2) or MDCK 33016, deposit number DSM ACC 2219 as described in WO 97/37001), cat, and rodent (e.g., hamster cells such as BHK21-F, HKCC cells, or Chinese hamster ovary cells (CHO cells)), and may be obtained from a wide variety of developmental stages, including for example, adult, neonatal, fetal, and embryo.

The polynucleotides encoding the SARS viral proteins may be modified to facilitate or enhance expression. For example, commercial leader sequences known in the art, such as tPA or IgK or interleukin-2, may be used in the recombinant constructs. Preferably, however, the natural SARS leader sequence is used. Use of the natural leader sequence can be used to ensure that the protein will be trafficked in human cells in the same way as during a normal viral infection, which may be advantageous e.g. for DNA vaccines, where antigen is expressed in situ.

As discussed above, tag sequences can be used in the expression constructs to facilitate purification, detection and stability of the expressed protein. Tag proteins suitable for use in the invention include a polyarginine tag (Arg-tag), polyhistidine tag (His-tag), FLAG-tag, Strep-tag, c-myc-tag, S-tag, calmodulin-binding peptide, cellulose-binding domain, SBP-tag,, chitin-binding domain, glutathione S-transferase-tag (GST), maltose-binding protein, transcription termination anti-termination factor (NusA), E. coli thioredoxin (TrxA) and protein disulfide isomerase I (DsbA). Preferred tag proteins include His-tag and GST. A full discussion on the use of tag proteins can be found at Terpe et al., "Overview of tag proteins in fusions: from molecular and biochemical fundamentals to commercial systems", Appl Microbiol Biotechnol (2003) 60:523-533.

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After purification, the tag proteins may optionally be removed from the expressed fusion protein, i.e., by specifically tailored enzymatic treatments known in the art. Commonly used proteases include enterokinase, tobacco etch virus (TEV), thrombin, and factor X.

One or more amino acid sequences or amino acid domains of the spike protein may be removed to facilitate mammalian recombinant expression. For instance, the entire S2 domain or the spike transmembrane region may be removed. Representative examples of some expression constructs of both full length and truncated spike glycoprotein suitable for mammalian expression are shown in FIGURE 40. Polynucleotide sequences representing each construct are shown in SEQ ID NOS 6578-6583. A description of each annotation is shown below:

Clone Name	Description	Expression Construct
nSh	natural leader sequence full length Spike	SEQ ID NO: 6578
	histidine tag	
nS	natural leader sequence full length Spike	SEQ ID NO: 6579
nSh∆TC	natural leader sequence	SEQ ID NO: 6580
	Spike without transmembrane sequence histidine tag	
nSΔTC	natural leader sequence	SEQ ID NO: 6581
nS1h	Spike without transmembrane sequence	
112111	natural leader sequence S1 domain	SEQ ID NO: 6582
;	histidine tag	
nS1	natural leader sequence	SEQ ID NO: 6583
* ja	S1 domain	

Cloned cDNA fragments that encompass full-length Spike coding sequences, as well as a Spike construct deleted of the transmembrane and cytoplasmic domains (TM-Cy-deleted Spike) for secretion were inserted into an expression vector pCMVIII to create nSh and nShΔTC, respectively. Both spike proteins were tagged with six histidine residues at the end of C-terminus to aid initial characterization of the expressed spike proteins. Similar sequences encoding full-length Spike or transmembrane and cytoplasmic domain deleted Spike, but without the histidine "tag" are readily substituted by one of skill in the art.

The likely locations of the expressed spike constructs was assessed by separating expressed proteins into an aqueous fraction (AF) and a detergent fraction (DF) using the procedure shown in Figure 48, with results of western blot analysis shown in Figure 43. The above described vector constructs were evaluated for expression after transfection into COS7 cells. The construct expressing the full length spike protein remained in the cell membrane while the construct expressing the truncated spike protein was located either in the cytosol (Figure 43) or secreted into the cell medium (Figure 44). As shown in Figure 43, full-length spike protein is found in DF (membrane) in an aggregated form, while the truncated protein is found in AF (cytosol) as a

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monomer. As shown in Figure 44, deleted proteins ($Sh\Delta TC$) are secreted, and a small fraction of full-length spike protein is detected in the medium by rabbit serum.

Recombinantly expressed spike proteins may be oligomerized. When the spike proteins are to be used in a vaccine or to generate antibodies specific to the spike protein, they are preferably oligomerized. In order to obtain oligomerized spike protein, it is preferred to maintain the transmembrane domain in the recombinant expression construct. For example, FIGURE 41 illustrates a western blot of COS7 cell lysates comparing expressed nSh and nSh Δ TC using both anti-his tag and rabbit anti-SARS antibodies. As shown full-length (nSh) aggregates, but the truncated (nSh Δ TC) spike protein does not. Antibody raised against the Histagged protein recognizes full-length and truncated spike proteins in native and reduced forms. Rabbit antiserum recognizes spike protein only in non-reducing conditions. Spike aggregates or oligomers were present in larger amounts in the cell lysates from the expressed nSh constructs. Preferably, the oligomerized spike proteins form a homotrimer, as indicated in FIGURE 47

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A further experiment, illustrated in FIGURE 42, demonstrates that the oligomerization of the expressed nSh constructs is likely due to a non-covalent linkage (and is likely not due to, for example, a disulfide bond). The oligomer dissociates into monomers at elevated temperature (80-100°C), but is stable in reducing conditions if not heated.

It is further preferred that recombinantly expressed spike proteins are glycoslyated. Tunicamycin and glycosidases were used to assess glycosylation. FIGURE 45 illustrates that glycoslation of expressed spike proteins is not affected by removal of the transmembrane domain region. Both full-length (Sh) and truncated (Sh Δ TC) SARS spike proteins are glycosylated.

Preferably, expression of the constructs of the invention is not toxic to the mammalian host cell. FIGURE 46 demonstrates that expression of the illustrated spike constructs is not toxic to the COS7 host cell.

Methods for transfecting, expressing, culturing, isolating and purifying recombinant proteins from mammalian cell cultures are known in the art. For example, the SARS spike constructs of the invention may be expressed in 293 cells. These cells may be cultured and transfected in static or monolayer cultures. For rapid large-scale production of SARS protein antigens in sufficient quantities for *in vitro* and *in vivo* evaluation, including immunogenicity studies, large-scale transient transfection of 293 (human embryonic kidney) cells may be used to obtain milligram quantities of the recombinant antigen(s). Alternatively, larger scale transfection of these cells may be performed with 293 cells in suspension culture. Preferably, the expressed SARS proteins are harvested from the transfected cells between 48 and 72 hours after transfection or even from 72 to 96 or more hours after transfection.

Where the host cells are transfected with truncated spike expression constructs, the expressed spike protein is secreted from the host cells and collected from the cell media. After

concentration, the spike protein may be purified from the media using, for example, GNA lectin followed by DEAE and ceramic hydroxyapatite column chromatography.

Where the host cells are transfected with full length spike expression constructs, but rather is retained within the cells, and may be purified from triton X-100 detergent extracted cells. The full-length Spike protein can then be captured on GNA lectin, followed by hydroxyapatite and SP chromatography.

Chinese Hamster Ovary (CHO) or other eukaryotic (e.g., mammalian) cells that stably express the SARS viral antigens of the invention may also be derived (e.g. Figure 73). Preferably, the cells are CHO cells, and these constructs will comprise one or more marker or selection genes in order to select for the desired CHO cells. In one embodiment, the constructs comprise a CMV enhancer/promoter, ampicillin resistance gene, and a fused DHFR and attenuated neomycin gene for selection purposes. Stable cell lines can then be produced using the neomycin selection system in CHOK-1 cells. Selected clones can then be sequenced to verify the integrity of the insert, and transient transfections can then be performed using Trans-LT1 polyamine transfection reagent (PanVera Corp., Madison, WI) to assess the expression level and also the integrity of the expressed protein by ELISA and western blot analysis.

Methods for derivation of CHO cells stably expressing the SARS viral antigens of the invention comprise the steps of transfection and primary screening with selective medium. Optionally, these steps are followed by subcloning to assure purity of cell lines. Cell culture supernatants can be assayed using an antigen capture ELISA to quantify expression levels at all stages of selection and amplification.

For full-length Spike expression constructs, methanol fixed cells can be screened for internal expression by immunofluorescent staining using a rabbit anti-SARS antibody. Successive measurements at the T75-flask stage of expansion can be employed to assure stability of expression levels. The molecular mass and integrity of the expressed proteins can be checked by PAGE both under native and reducing and denaturing conditions, followed by immunoprobing.

In one embodiment, the pCMV3 vectors expressing SARS-CoV Spike proteins in either full-length or truncated forms is introduced into CHOK-1 cells using the Trans-LT-1 reagent. On day one, 1x10⁶ cells are plated on 100 mm dishes in non-selective F12 media + 10% Fetal Bovine Serum + 4 mM Glutamine. On day two, the cells are transfected with a DNA:LT-1 mixture and the media then replaced with complete F12 media. Twenty-four to forty-eight hours later depending on the cell density, each 100 mm dish is split to 4-6 100 mm dishes. The medium is changed to complete selective media containing Geneticin (neomycin) at 500 µg/ml. All bovine serum used in these procedures is from TSE-free sources that meet current FDA standards. Twenty-four hours later the medium is changed to complete selective medium plus

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500 ug/ml neomycin. Ten to fourteen days later, individual colonies are picked and transferred to 96 well plates and cultured in complete selective medium but without G418. When approximately 80% of the wells are confluent, twenty-four hour supernatants are screened by spike capture ELISA positive clones are transferred to twenty-four well plates. For the initial expression of full length Spike protein, methanol fixed cells will be screened by immunoflourescent staining using a rabbit anti-SARS antibody. After the low expressing cell lines have been eliminated and there are less than 20-30 cell lines, capture ELISA and westerns will be used to determine the expression level after cell lysis. A portion of each cell line will be pelleted, weighed and lysed in 1% triton lysis buffer containing MOPS, NaCl and MgCl₂ at the same ratio of cell weight to lysis buffer. After lysis the supernatant is collected and expression level is determined. Three to four clones producing the highest levels of spike protein in correct structure and conformation will be grown in three-liter bioreactors for expansion and adaptation to low serum suspension culture conditions for scale-up.

The antigen capture ELISA assay for the SARS spike protein can be performed as described in the art. A brief description of this assay follows. 96 well flat-bottom plates (Corning, Corning, NY) are coated with 250ng per well of purified immunoglobulin obtained from rabbit sera that were immunized with inactivated SARS virus. Between steps, the plates are washed in a buffer containing 16%NaCl and 1% Triton X100. 100µL of supernatant or lysate samples (diluted in a buffer containing 100mM NaPO₄, 0.1% Casein, 1mM EDTA, 1% Triton X100, 0.5M NaCl and 0.01% Thiomersal, pH 7.5) are added and incubated for 2 hours at 37°C. Bound antigen is reacted against pooled SARS+ve serum or high affinity monoclonal antibody either human or mouse against SARS spike protein (1 hour incubation, 37°C) and detected using appropriate species-specific peroxidase conjugated second antibody (30 minute incubation at 37°C; TAGO, Burlingame, CA). The plates are developed for 15 minutes at room temperature using TMB substrate (Pierce, Rockford, IL) and the reaction stopped using 4N phosphoric acid. The plates are read at a wavelength of 450nm and the concentration of protein per ml sample is derived from a standard curve (OD vs. protein concentration) based on serial dilutions of a known concentration of recombinant spike protein.

The immunoprobing analysis can also be performed following the standard methods described elsewhere in the art. A brief description follows. 10-20 µl of the sample is analyzed on 4-20% SDS PAGE under non-reducing/ denaturing conditions with mild heating. The gels are run for 1.5-2.0 hours at 100V constant voltage. The proteins are then transferred onto nitrocellulose membranes (Millipore, Bedford, MA) for 45 min using the semidry western transfer system (BioRad, Hercules, CA) following the manufacturer's instructions. The membrane is then reacted against polyclonal anti-spike rabbit serum, followed by anti-rabbit Ig

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conjugated to Alexa 688 (Molecular Probes, Oregon). The blots are scanned using an infrared imaging system (LI-Cor, Inc., Lincoln, Nebraska).

The highest expressing candidate cell lines can be screened for spike protein expression and stability in small-scale (3 liter) suspension cultures. The candidate clone can be further evaluated for level of expression as well as integrity of expressed protein after amplification, and subsequently tested for expression stability in the absence of selection. The selected clones can also be tested for maintenance of the DNA sequence integrity of the integrated SARS spike protein gene. To quickly monitor the expression levels in small flask (T25 or T75) and in the three liter evaluation cultures, a lectin-based process (Gluvanthus Nivalis lectin) may be used to isolate SARS spike protein to a degree of purity that allows semi-quantitation and characterization of the protein in CHO supernatant. For full-length spike protein, it will be obtained from triton X-100 detergent extracted cells. Full-length Spike protein will be then captured on GNA lectin, followed by hydroxyapatite and SP chromatograph. Eluted protein is then characterized by: 1) polyacrylamide gel electrophoresis (PAGE) and Coomassie staining, 2) Immunoprobing with anti-SARS rabbit sera, 3) structural characterization using size exclusion chromatography (SEC), as well as mass spec analysis using MALDI-TOF.

Routes and methods of immunization of the vaccines of the invention are discussed in more detail in a section below. Examples 7 to 9 illustrate sample immunization protocols for the recombinant spike proteins.

Vaccine testing

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Prior to human administration, it is normal to test vaccines in animal models. A mouse model of SARS coronavirus infection is known (Subbarao et al. (2004) J Virol 78:3572-77), and other animals that may be used as models of infection and/or disease include ferrets and monkeys. Thus the invention provides a non-human animal that is infected by the SARS coronavirus, wherein the animal is preferably a ferret or a primate (e.g. a monkey or a macaque). The animal may be gnotobiotic. The animal is preferably not a cat (Felis domesticus). The animal may or may not display SARS disease symptoms e.g. ferrets (Mustela furo) show prominent pulmonary pathology after infection. See: Martina et al. (2003) Nature 425:915. E. Polynucleotides encoding the SARS Antigens of the Invention

The invention includes polynucleotides encoding for the SARS antigens of the invention. In addition, the invention includes polynucleotides which have been optimized for recombinant production (e.g. codon optimization) of the SARS antigens of the invention, including polynucleotides encoding for each of the SARS fusion constructs discussed above.

F. Viral vector or Viral Particle delivery of the SARS Antigens of the Invention

The antigens of the invention may be expressed in vivo or in vitro by polynucleotides encoding the antigens. Expression and delivery of the polynucleotides of the invention may be facilitated via viral vectors and/or viral particles.

Gene-based delivery systems derived from viruses, such as alphaviruses, are useful for the ex vivo and in vivo administration of heterologous genes, including one or more SARS genes, having therapeutic or prophylactic applications. These systems can also be used for the production of recombinant proteins derived from the SARS virus in cultured cells. Gene-based delivery systems of the invention include viral vectors (e.g., adenovirus vector, poxvirus vector, alphavirus vector) and non-viral nucleic acid vectors (e.g., DNA, RNA) encoding one or more SARS virus antigens. Polynucleotides encoding SARS virus antigen(s) are incorporated into the gene-based vaccines individually or in combination (e.g., as bicistronic constructs).

1. Alphavirus

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Alphaviruses are members of *Togaviridae* family and share common structural and replicative properties. Sindbis virus (SIN) is the prototype virus for the molecular study of other alphaviruses, and together with Venezuelan equine encephalitis virus (VEE) and Semliki Forest virus (SFV), are the most widely utilized alphaviruses being developed into expression vectors for heterologous genes (Schlesinger and Dubensky (1999) *Curr Opin. Biotechnol. 10*:434-439; Schlesinger (2001) Expert Opin. Biol. Ther. 1:177-91).

Alphaviruses possess a relatively small single-stranded RNA genome of positive polarity, which is approximately 12 kb in length, capped and polyadenylated. The RNA interacts with viral capsid protein monomers to form nucleocapsids, which in turn, are surrounded by a host cell-derived lipid envelope from which two viral glycoproteins, E1 and E2, protrude forming "spike" trimers of heterodimeric subunits. Two open reading frames (ORFs) encode as polyproteins the enzymatic nonstructural replicase proteins (5' ORF) and the virion structural proteins (3' ORF). The structural polyprotein is translated from a highly abundant subgenomic mRNA, which is transcribed from a strong internal alphavirus promoter (Strauss and Strauss (1994) Microbiol. Rev. 58:491-562). Replication of the genome occurs exclusively within the host cell cytoplasm as RNA.

The most common alphavirus expression vectors have exploited both the positive-stranded nature and modular organization of the RNA genome. These vectors, termed "replicons" due to their property of self-amplification, permit insertion of heterologous sequences in place of the structural polyprotein genes, while maintaining the 5'- and 3'-end cis replication signals, the nonstructural replicase genes, and the subgenomic junction region promoter (Xiong et al. (1989) Science 243:1188-1191; Liljestrom (1991) Bio/Technology 9:1356-1361). Chimeric alphavirus vectors (and particles) from sequences derived from divergent virus families have also been -140-

described. (see, for example United States patent application serial number 09/236,140; see also, US Patents 5,789,245, 5,842,723, 5,789,245, 5,842,723, and 6,015,694; as well as WO 95/07994, WO 97/38087 and WO 99/18226). Co-owned International Publication WO 02/099035, published December 12, 2002 and incorporated by reference in its entirety herein, describes chimeric alphavirus molecules and modified alphavirus molecules having modified Biosafety Levels.

The absence of structural protein genes renders alphavirus replicon vectors defective, in that RNA amplification and high-level heterologous gene expression occurs within the target cell, but cell-to-cell spread of vector is not possible due to the inability to form progeny virions. Through the years, several synonymous terms have emerged that are used to describe alphavirus replicon particles. These terms include recombinant viral particle, recombinant alphavirus particle, alphavirus replicon particle and replicon particle. However, as used herein, these terms all refer to a virion-like unit containing an alphavirus-derived RNA vector replicon. Moreover, these terms may be referred to collectively as vectors, vector constructs or gene delivery vectors.

Packaging of replicon RNA into particles can be accomplished by introducing the replicon RNA into permissive cells (e.g., RNA or DNA transfection, or particle infection) that also contain one or more structural protein expression cassettes or "defective helper" constructs encoding the alphavirus structural proteins. These structural protein encoding constructs may themselves be introduced into the cells by transfection of either RNA or DNA, and most commonly retain the native alphavirus subgenomic promoter, as well as 5'- and 3'-end cis signals for co-amplification with the replicon, but are devoid of any replicase genes and the RNA packaging signal (Liljestrom (1991) Bio/Technology 9:1356-1361; Pushko et al. (1997) Virology 239:389-401; Polo et al. (1999) PNAS 96:4598-4603). Permanent cell lines that are stable transformed with constructs expressing the alphavirus structural proteins (e.g., packaging cell lines) offer a means to avoid transient transfection production methods (Polo et al. (1999) PNAS 96:4598-4603).

The present invention includes compositions and methods for the production of replication defective viral vector particles (e.g., alphavirus replicon particles) for use in the ex vivo and in vivo administration of heterologous genes encoding proteins having therapeutic or prophylactic application, including genes encoding for one or more SARS viral antigens.

In one aspect, the invention includes a method of producing replication defective viral vector particles (e.g., alphavirus replicon particles) comprising the steps of introducing at least one nucleic acid molecule comprising a viral vector (e.g., alphavirus replicon RNA) into immortalized cells of the present invention, under conditions that allow for complementation of the viral vector (e.g., alphavirus replicon RNA) and production of viral vector particles (e.g., alphavirus replicon particles), and isolating the viral vector particles from the cells or cell culture

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supernatants. In certain embodiments, the immortalized cells are grown in suspension, for example PERC.6 cells. In other embodiments, the methods are performed in large-scale volumes, for example, liter volumes or greater, such as for example in roller bottles, large flasks. Nunc Cell Factories, Corning Cell Cubes, fermentation vessels, etc).

In certain embodiments, the viral vector is an alphavirus replicon RNA that requires complementation by providing one or more alphavirus structural proteins in trans, within the immortalized cell. In such instances, the methods of complementation to produce alphavirus replicon particles may involve the introduction of one or more nucleic acids (e.g., RNA, DNA) encoding said alphavirus structural protein(s) (e.g., capsid and/or envelope glycoproteins) into the immortalized cells, either transiently or stably, and either concurrent with or prior to the introduction of the alphavirus replicon RNA. In certain embodiments, the alphavirus replicon RNA is introduced into the cell by transfection an in vitro transcribed RNA. In other embodiments, the alphavirus replicon RNA is introduced into the cell by transfection of a DNA (e.g., ELVIS), which is capable of transcribing within the cell, the replicon RNA. In yet other embodiments, the alphavirus replicon RNA is introduced into the cell by infection with a seed stock of alphavirus replicon particles. In certain embodiments, the nucleic acids encoding said alphavirus structural protein(s) are defective helper RNA or are DNA that can transcribe within the cell defective helper RNAs.

As discussed herein, "alphavirus RNA replicon vector", "RNA replicon vector", "replicon :0 vector" or "replicon" refers to an RNA molecule that is capable of directing its own amplification or self-replication in vivo, within a target cell. To direct its own amplification, the RNA molecule should encode the polymerase(s) necessary to catalyze RNA amplification (e.g., alphavirus nonstructural proteins nsP1, nsP2, nsP3, nsP4) and also contain cis RNA sequences required for replication which are recognized and utilized by the encoded polymerase(s). An alphavirus RNA vector replicon should contain the following ordered elements: 5' viral or cellular sequences required for nonstructural protein-mediated amplification (may also be referred to as 5' CSE, or 5' cis replication sequence, or 5' viral sequences required in cis for replication, or 5' sequence which is capable of initiating transcription of an alphavirus), sequences which, when expressed, code for biologically active alphavirus nonstructural proteins (e.g., nsP1, nsP2, nsP3, nsP4), and 3' viral or cellular sequences required for nonstructural protein-mediated amplification (may also be referred as 3' CSE, or 3' viral sequences required in cis for replication, or an alphavirus RNA polymerase recognition sequence). The alphavirus RNA vector replicon also should contain a means to express one or more heterologous sequence(s), such as for example, an IRES or a viral (e.g., alphaviral) subgenomic promoter (e.g., junction region promoter) which may, in certain embodiments, be modified in order to increase or reduce viral transcription of the subgenomic fragment, or to decrease homology with

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defective helper or structural protein expression cassettes, and one or more heterologous sequence(s) to be expressed. Preferably the heterologous sequence(s) comprises a protein-encoding gene, which is the 3' proximal gene within the vector replicon. And preferably the replicon further comprises a polyadenylate tract.

As discussed herein, "recombinant Alphavirus Particle", "alphavirus replicon particle" and "replicon particle" refers to a virion-like unit containing an alphavirus RNA vector replicon. Generally, the recombinant alphavirus particle comprises one or more alphavirus structural proteins, a lipid envelope and an RNA vector replicon. Preferably, the recombinant alphavirus particle contains a nucleocapsid structure that is contained within a host cell-derived lipid bilayer, such as a plasma membrane, in which one or more alphaviral envelope glycoproteins (e.g., E2, E1) are embedded. The particle may also contain other components (e.g., targeting elements such as biotin, other viral structural proteins or portions thereof, hybrid envelopes, or other receptor binding ligands), which direct the tropism of the particle from which the alphavirus was derived. Generally the interaction between alphavirus RNA and structural protein(s) necessary to efficiently form a replicon particle or nucleocapsid may be an RNA-protein interaction between a capsid protein and a packaging signal or packaging sequence contained within the RNA.

"Alphavirus packaging cell line" refers to a cell which contains one or more alphavirus structural protein expression cassettes and which produces recombinant alphavirus particles (replicon particles) after introduction of an alphavirus RNA vector replicon, eukaryotic layered vector initiation system, or recombinant alphavirus particle. The parental cell may be of mammalian or non-mammalian origin. Within preferred embodiments, the packaging cell line is stably transformed with the structural protein expression cassette(s).

"Defective helper RNA" refers to an RNA molecule that is capable of being amplified and expressing one or more alphavirus structural proteins within a eukaryotic cell, when that cell also contains functional alphavirus nonstructural "replicase" proteins. The alphavirus nonstructural proteins may be expressed within the cell by an alphavirus RNA replicon vector or other means. To permit amplification and structural protein expression, mediated by alphavirus nonstructural proteins, the defective helper RNA molecule should contain 5'-end and 3'-end RNA sequences required for amplification, which are recognized and utilized by the nonstructural proteins, as well as a means to express one or more alphavirus structural proteins. Thus, an alphavirus defective helper RNA should contain the following ordered elements: 5' viral or cellular sequences required for RNA amplification by alphavirus nonstructural proteins (also referred to elsewhere as 5' CSE, or 5' cis replication sequence, or 5' viral sequences required in cis for replication, or 5' sequence which is capable of initiating transcription of an alphavirus), a means to express one or more alphavirus structural proteins, gene sequence(s) which, when expressed,

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codes for one or more alphavirus structural proteins (e.g., C, E2, E1), 3' viral or cellular sequences required for amplification by alphavirus nonstructural proteins (also referred to as 3' CSE, or 3' viral sequences required in cis for replication, or an alphavirus RNA polymerase recognition sequence), and a preferably a polyadenylate tract. Generally, the defective helper RNA should not itself encode or express in their entirety all four alphavirus nonstructural proteins (nsP1, nsP2, nsP3, nsP4), but may encode or express a subset of these proteins or portions thereof, or contain sequence(s) derived from one or more nonstructural protein genes. but which by the nature of their inclusion in the defective helper do not express nonstructural protein(s) or portions thereof. As a means to express alphavirus structural protein(s), the defective helper RNA may contain a viral (e.g., alphaviral) subgenomic promoter which may, in certain embodiments, be modified to modulate transcription of the subgenomic fragment, or to decrease homology with replicon RNA, or alternatively some other means to effect expression of the alphavirus structural protein (e.g., internal ribosome entry site, ribosomal readthrough element). Preferably an alphavirus structural protein gene is the 3' proximal gene within the defective helper. In addition, it is also preferable that the defective helper RNA does not contain sequences that facilitate RNA-protein interactions with alphavirus structural protein(s) and packaging into nucleocapsids, virion-like particles or alphavirus replicon particles. A defective helper RNA is one specific embodiment of an alphavirus structural protein expression cassette.

Alphavirus for use in the invention may be grown in any one of the cell lines discussed above as suitable for the SARS virus.

Alphavirus replicon particles may be produced according to the present invention by using the above cell lines (e.g., immortalized cell lines) and a variety of published and accepted alphavirus vector methodologies. Such methodologies include, for example, transient packaging approaches, such as the co-transfection of in vitro transcribed replicon and defective helper RNA(s) (Liljestrom, Bio/Technology 9:1356-1361, 1991; Bredenbeck et al., J. Virol. 67:6439-6446, 1993; Frolov et al., J. Virol. 71:2819-2829, 1997; Pushko et al., Virology 239:389-401, 1997; US Patents 5,789,245 and 5,842,723) or co-transfection of plasmid DNA-based replicon and defective helper construct(s) (Dubensky et al., J. Virol. 70:508-519, 1996), as well as introduction of alphavirus structural protein expression cassettes (e.g., DNA-based defective helper) into immortalized cell lines of the present invention to create stable packaging cell lines (PCL) (Polo et al., PNAS 96:4598-4603, 1999; US Patents 5,789,245, 5,842,723, 6,015,694; WO 97/38087, WO 99/18226, WO 00/61772, and WO 00/39318). Stable packaging cell lines may then be utilized for alphavirus replicon particle production. The PCL may be transfected with in vitro transcribed alphavirus replicon RNA, transfected with a plasmid DNA-based replicon (e.g., ELVIS vector), or infected with a seed stock of alphavirus replicon particles, and then incubated under conditions and for a time sufficient to produce progeny alphavirus replicon particles in the

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culture supernatant. In addition, progeny replicon particles can subsequently be passaged in additional cultures of naïve PCL by infection, resulting in further expansion and commercial scale preparations. Importantly, by using defective helper RNA or stable PCL based on the "split" structural gene configuration, these replicon particle stocks may be produced free from detectable contaminating RCV.

Following harvest, crude culture supernatants containing the chimeric alphavirus replicon particles may be clarified by passing the harvest through a filter (e.g., 0.2 uM, 0.45 uM, 0.65 uM, 0.8 uM pore size). Optionally, the crude supernatants may be subjected to low speed centrifugation prior to filtration to remove large cell debris. Within one embodiment, an endonuclease (e.g., Benzonase, Sigma #E8263) is added to the preparation of alphavirus replicon particles before or after a chromatographic purification step to digest exogenous nucleic acid. Further, the preparation may be concentrated prior to purification using one of any widely known methods (e.g., tangential flow filtration). Crude or clarified alphavirus replicon particles may be concentrated and purified by chromatographic techniques (e.g., ion exchange chromatography, size exclusion chromatography, hydrophobic interaction chromatography, affinity chromatography), such as those described in WO01/92552, incorporated by reference in its entirety herein. Two or more such purification methods may be performed sequentially.

EXAMPLE OF ALPHAVIRUS REPLICON PARTICLES ENCODING SARS VIRUS SPIKE (S)
ANTIGEN

The invention includes compositions and methods for the production of replication defective viral vector particles (e.g., alphavirus replicon particles) for use in the ex vivo and in vivo administration of heterologous genes encoding proteins having therapeutic or prophylactic application, including genes encoding for one or more SARS viral antigens.

The following example illustrates a method of preparing alphavirus replicon particles encoding SARS virus spike (s) antigen.

The SARS virus spike gene can be incorporated into alphavirus replicon particles derived from a variety of alphavirus, such as Sindbis virus, Semliki Forest virus (US 5739026), Venezuelan equine encephalitis virus (US 6531135), and replicon particle chimeras derived from more than one alphavirus (US 6376236, WO 02/99035). In addition, the SARS virus spike gene can be incorporated in its entirety (encoding full-length spike protein) or in a modified form that includes, for example, sequence deletions or truncations, such that the encoded a spike protein is of less than full-length (e.g., C-terminal truncation of one or more (e.g. at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25, 30 etc.) amino acids, deleted of transmembrane region and cytoplasmic tail).

For example, the spike gene may be cloned as a full-length gene into the VCR-chim2.1 vector (WO 02/99035) by standard RT-PCR conditions or by standard subcloning from one of the other plasmids described herein, using commercially available restriction endonucleases. For

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the reverse transcription step in standard RT-PCR, the Superscript pre-amplification kit (InvitrogenTM) and the primer SEQ ID NO: 7325 (sp-RT-R) are used:

For the amplification step, the cDNA polymerase advantage kit (Clonetech) and two primers Sp-F-BbvCI (SEQ ID NO: 7326) and Sp-R-NotI (SEQ ID NO: 7327) are used:

The forward primer is designed to contain the ccacc sequence (Kozak, 1991 *JBC* 19867-70) in front of the ATG codon to optimize translation efficiency of the spike gene. Also, the forward primer contains the BbvCI restriction site and the reverse primer contains the NotI restriction site for subsequent cloning of the PCR amplified gene.

The PCR product is purified using the QIAquick Nucleotide Removal kit (QIAgen), digested with BbvCl and Notl, gel purified with QIAquick Gel Extraction kit (QIAgen), and ligated to plasmid VCR-Chim2.1 pre-digested with the same enzymes. Clones containing the SARS spike sequence are verified by sequencing and the new construct is called VCR-Chim2.1-SARSspike.

To generate VEErep/SINenv-SARSspike replicon particles the plasmids VCR-Chim2.1-SARSspike, VCR-DH-Scap (WO 02/99035), and VCR-DH-Sglydl160 (WO 02/99035) are linearized with the restriction enzyme PmeI and used for *in vitro* transcription as described previously (Polo *et al.* 1999, PNAS 96: 4598-603; WO02/99035). The transcripts are cotransfected into BHK cells as previously described (Polo *et al.*, 1999, *ibid.*; WO02/99035). The transfected cells are incubated at 34 °C, the supernatants collected at 20 and 30 hrs postelectroporation, clarified by centrifugation, and purified by chromatography as previously described (WO 01/92552).

Expression of the SARS spike protein from the replicon particle vector is verified by infecting BHK cells overnight with purified VEErep/SINenv-SARSspike or VEErep/SINenv-GFP (WO 02/99035) replicon particles. In addition, BHK cells also were transfected in parallel with in vitro transcribed VCR-Chim2.1-SARSspike replicon RNA. At 16 hrs post-infection and transfection cells are lysed and a sample of the lysate analyzed by western blot using an antibody that recognizes SARS virus spike protein. The proteins on the gel are stained or transferred to a membrane for Western blot analysis with sera from convalescent patients or alternatively murine or rabbit antisera generated against SARS virus. VEErep/SINenv-SARSspike replicon particles are administered to the vaccine recipient (e.g., rodent, non-human primate, human) as described elsewhere in the present invention.

Figure 67 shows data from western blot analysis performed under non-reducing conditions, using a SARS virus specific rabbit polyclonal antisera. The western data demonstrate that not only is SARS spike protein expressed in cells infected with alphavirus replicon particles or transfected with replicon RNA, but the predominant form of spike is that of a homotrimer (Fig.67A). Similar homotrimeric association of the spike protein was observed in western blots

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of SARS virions purified from SARS virus infected VERO cell supernatants, and this homotrimer is heat labile, as indicated by the dissociation into monomeric forms at 80°C and 100°C (Fig.67B).

To further characterize SARS Spike protein expression and processing following expression from alphavirus replicon vectors, BHK-21 cells were infected with alphavirus replicon particles expressing the full-length Spike. At 6 hr post-infection with an MOI of 5, infected cells were labeled for 1 hr with L-[35] methionine/cysteine and chased for the indicated time. The [35]-labeled spike protein was immunoprecipitated by anti-SARS rabbit serum and digested with Endo-H. Both digested and undigested proteins were analysed by 4% polyacrylamide-SDS PAGE under reducing conditions. As shown in Figure 55, the full-length spike protein is synthesized as an Endo-H sensitive high mannose glycoprotein (gp170, an ER form) that undergoes modification to an Endo-H resistant glycoprotein with complex oligosaccharides (gp180, a Golgi form). The conversion of gp170 into the gp180 form takes place within 2 hr.

Alphavirus replicon particles expressing one or more SARS proteins (e.g., VEErep/SINenv-SARSspike replicon particles) are administered to the vaccine recipient in order to induce a SARS specific immune response (e.g., rodent, ferret, non-human primate, human) as described elsewhere in the present invention. Immunization may be performed through a variety of routes, including for example, intramuscular, subcutaneous, intradermal, and intranasal. In additon, the alphavirus replicon particles may be used alone or in combination (e.g., "primeboost") with other vaccine approaches of the present invention, or alternatively the alphavirus replicon particles may co-express antigen from other respiratory pathogens or be co-administered in combination with alphavirus replicon particles expressing antigens from other respiratory pathogens (e.g., influenza virus, parainfluenza virus, respiratory syncytial virus, human metapneumovirus). For example, the induction of anti-spike protein antibodies in animals immunized IM with VEErep/SINenv-SARSspike replicon particles was demonstrated in mice (Figure 68). These mouse studies also included additional vaccine groups for comparison, including the inactivated SARS virus and recombinant truncated spike protein vaccines describe elsewhere herein, as well as plasmid DNA used as a prime, followed by alphavirus replicon particles as a boost. The data clearly show very potent immune responses for all vaccine groups, including the alphavirus replicon particle group. It should be noted that the level of antibody induced by the inactivated SARS virus vaccine used in these experiments has been shown to be protective in a SARS virus animal challenge model.

Similarly, genes encoding other SARS virus antigens (e.g., nucleocapsid protein, membrane glycoprotein) are cloned into alphavirus replicon vectors, either individually or in

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combination, to generate alphavirus replicon particles according to the teachings of the present invention and using standard molecular biology techniques..

EXAMPLE OF ALPHAVIRUS-BASED PLASMID DNA EXPRESSING SARS VIRUS SPIKE (S)

The invention includes preparation of plasmid DNA expressing a SARS virus antigen for prophylactic or therapeutic immunization against SARS virus infection. In one embodiment, the SARS viral antigen is a spike (S) protein. In one embodiment, the plasmid DNA is alphavirus-based.

The following example illustrates one method for preparing an alphavirus-based plasmid

DNA expressing SARS virus spike (S).

SARS spike gene can be delivered using any of the alphavirus-based plasmid DNA replicons such as ELVS (Dubensky et al, 1996 J Virol. 70: 508-19), SINCP (WO 01/81609), or VCP (PCT WO 02/99035).

For example, the SARS spike gene is cloned into SINCP using the standard RT-PCR techniques. The oligo Sp-RT-R is used for the reverse transcription step with the Superscript preamplification kit (Invitrogen). For the amplification step, the cDNA polymerase advantage kit (Clonetech) with the Sp-R-NotI and Sp-F-XhoI (SEQ ID NO: 7328) primers is used.

The Sp-F-XhoI primer was designed to contain the ccacc sequence in front of the ATG codon to optimize translation efficiency (Kozak 1991, ibid) of the spike gene. Also, the primer contains the XhoI restriction site for the subsequent cloning of the PCR amplified gene.

The PCR product is purified using the QIAquick Nucleotide removal kit, digested with XhoI and NotI, gel purified with QIAquick Gel Extraction kit, and ligated to plasmid SINCP predigested with the same enzymes. Clones containing the SARS spike sequence are verified by sequencing and the new construct is called SINCP-SARS spike.

Expression of the SARS spike gene is verified by transient transfection of BHK cells with $2\mu g$ of either plasmid DNA SINCP-SARSspike or SINCP pre-incubated for 5 minutes with 5 μl of TransIT Polyamine reagent (Mirrus) in low serum medium Optimem (Life Technologies). At 48 hrs pos-transfection cells are lysed and a sample of the lysate is run on 8% SDS-PAGE. The proteins on the gel are either stained or transferred to a membrane for Western blot analysis with sera from convalescent patients, or alternatively with sera from mouse or rabbits.

SINCP-SARSspike plasmid replicon is administered to the vaccine recipient (e.g., rodent, non-human primate, human) as a formulated or unformulated plasmid vaccine, alone or in combination (e.g., "prime-boost") with other vaccines of the present invention, as described elsewhere herein.

Similarly, genes encoding other SARS virus antigens (e.g., nucleocapsid protein, membrane glycoprotein) are cloned into alphavirus plasmid replicon vectors.

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2. Plasmid Expression Vectors

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EXAMPLE OF PLASMID DNA EXPRESSING SARS VIRUS SPIKE (S)

The following example illustrates a method for preparing plasmid DNA expressing SARS virus spike (s).

The SARS virus spike antigen also may be delivered using other plasmid DNA expression vectors (sometimes referred to as "conventional" DNA vaccines), based on a polymerase II promoter, such as, for example, a CMV promoter. A DNA vaccine of the spike antigen gene induces an antibody response in mice (Zhao et al. (2004) Acta Biochim et Biophysica Sinica 36:37-41), and has been found to induce viral neutralization and protective immunity in mice (Yang et al. (2004) Nature 428:561-564), particularly when truncated at the C-terminus.

For example, the SARS spike gene is cloned into pCMVKm2 (Zur Megede et al., J. Virol., 74:2628-2635, 2000; SEQ ID NO: 9923) using standard RT-PCR techniques. The oligo Sp-RT-R is used for the reverse transcription step with the Superscript pre-amplification kit (Invitrogen). For the amplification step, the cDNA polymerase advantage kit (Clonetech) is used with primers Sp-F-EcoRI (SEQ ID NO: 7329) and Sp-R-XbaI (SEQ ID NO: 7330).

The forward primer was designed to contain the CCACC (SEQ ID NO: 7331) sequence in front of the ATG codon to optimize translation efficiency (Kozak 1991, ibid.) of the spike gene. Also, the forward primer contains the EcoRI restriction site and the reverse primer contains the XbaI restriction site for the subsequent cloning of the PCR amplified gene.

The PCR product is purified using the QIAquick Nucleotide Removal kit, digested with XhoI and NotI, gel purified with QIAquick Gel Extraction kit, and ligated to plasmid pCMVKm2 pre-digested with the same enzymes. Clones containing the SARS spike sequence are verified by sequencing and the new construct is called pCMVKm2-SARSspike.

Expression of the SARS spike gene is verified by transient transfection of BHK or 293 cells with $2\mu g$ of either plasmid DNA pCMVKm2-SARSspike or pCMVKm2 pre-incubated for 5 minutes with $5\,\mu l$ of TransIT Polyamine reagent (Mirrus) in low serum medium Optimem (Life Technologies). At 48 hrs pos-transfection cells are lysed and a sample of the lysate is run on 8 % SDS-PAGE. The proteins on the gel are either stained or transferred to a membrane for Western blot analysis with sera from convalescent patients, or alternatively using mouse or rabbit antisera

Plasmid pCMVKm2-SARSspike is administered to the vaccine recipient (e.g., rodent, nonhuman primate, human) as a formulated or unformulated plasmid vaccine, as described elsewhere in the present invention.

Similarly, genes encoding other SARS virus antigens (e.g., nucleocapsid protein, membrane glycoprotein) are cloned into plasmid expression vectors

3. Virus-Like Particles comprising SARS antigens

The SARS viral antigens of the invention may be formulated into Virus Like Particles ("VLPs"). The invention thus includes virus-like particles (or VLPs) comprising one or more SARS viral antigens. Preferably, the VLPs comprise one or more SARS viral antigens selected from the group consisting of Spike (S), nucleocapsid (N), membrane (M) and envelope (E). Preferably, the VLPs comprise at least M and E.

The VLPs of the invention comprise at least one particle-forming polypeptide. Said particle-forming polypeptide is preferably selected from a Coronavirus structural protein. In one embodiment, the particle-forming polypeptide is selected from one or more SARS viral antigens. In another embodiment, the particle-forming polypeptide is selected from the structural protein of a non-SARS Coronavirus, such as, for example, Mouse Hepatitis Virus.

VLPs can be formed when viral structural proteins are expressed in eukaryotic or prokaryotic expression systems. Upon expression, the structural proteins self-assemble to form particles. Alternatively, viral structural proteins may be isolated from whole virus and formulated with phospholipids. Such viral structural proteins are referred to herein as "particle-forming polypeptides". VLPs are not infectious because no viral genome is present, however, these non-replicating, virus capsids mimic the structure of native virions.

Due to their structure, VLPs can display a large number of antigenic sites on their surface (similar to a native virus). VLPs offer an advantage to live or attenuated vaccines in that they are much safer to both produce and administer, since they are not infectious. VLPs have been shown to induce both neutralizing antibodies as well as T-cell responses and can be presented by both class I and II MHC pathways.

Previous work creating VLPs from coronavirus indicates that E and M proteins along may be sufficient for coronavirus VLP formation. See Fischer et al., J. Virol. (1998) 72:7885-7894 and Vennema et al. EMBO J. (1996) 15:2020-2028.

Chimeric VLPs comprising particle-forming polypeptides or portions thereof from non-SARS Coronaviruses are also included in the invention. Such particle-forming polypeptides may comprise a full length polypeptide from a non-SARS Coronavirus. Alternatively, a particleforming fragment may be used.

In one embodiment, a fragment of a non-SARS particle-forming polypeptide and a fragment of a SARS viral antigen are fused together. For instance, such chimeric polypeptides may comprise the the endodomain and transmembrane domain of a non-SARS particle-forming polypeptide and the ectodomain of a SARS viral antigen. In one example, the VLPs of the invention comprise a chimeric spike protein comprising an endodomain and transmembrane domain of the spike protein of Mouse Hepatitis Virus (MHV) and the chimeric spike protein further comprises the ectodomain of the SARS spike protein. Such VLPs may further comprise

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Coronavirus M and E proteins. Said M and E proteins may be selected from any coronavirus, including Mouse Hepatitis Virus (MHV) or SARS. Sample sequences of S, M and E proteins of MHV are included in the figures, *supra*.

Chimeric spike proteins derived from the ectodomain of feline infectious peritonitis virus (FIPV) spike protein fused to the endo and transmembrane domains of MHV spike protein have been previously disclosed. See WO 98/49195 and WO 02/092827. In these chimeric VLP structures, the capsid structure of the VLPs is formed by the M and E protein of MHV. The chimeric spike protein provides for the surface exposure of the ectodomain of the FIPV spike protein.

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As used herein, the term "virus-like particle" or "VLP" refers to a non-replicating, empty virus shell. VLPs are generally composed of one or more viral proteins, such as, but not limited to those proteins referred to as capsid, coat, shell, surface and/or envelope proteins, or particle-forming polypeptides derived from these proteins. VLPs can form spontaneously upon recombinant expression of the protein in an approrpirate expression system. Alternatively, viral structural proteins may be isolated from whole virus and formulated with phospholipids. Methods for producing particular VLPs are known in the art and discussed more fully below. The presence of VLPs in a composition can be detected using conventional techniques known in the art, such as by electron microscopy, x-ray crystallography, and the like. See, e.g., Baker et al., Biophys. J. (1991) 60:1445-1456; Hagensee et al., J. Virol. (1994) 68:4503-4505. For example, cryoelectron microscopy can be performed on vitrified aqueous samples of the VLP preparation in question, and images recorded under appropriate exposure conditions.

The phrase "particle-forming polypeptide" includes a full-length or near full-length viral protein, as well as a fragment thereof, or a viral protein with internal deletion, which has the ability to form VLPs under conditions that favor VLP formation. Accordingly, the polypeptide may comprise the full-length sequence, fragments, truncated and partial sequences, as well as analogs and precursor forms of the reference molecule. The term therefore includes deletions, additions and substitutions to the sequence, so long as the polypeptide retains the ability to form a VLP. Thus, the term includes natural variations of the specified polypeptide since variations in coat proteins often occur between viral isolates. The term also includes deletions, addition and substitutions that do not naturally occur in the reference protein, so long as the protein retains the ability to form a VLP.

Preferred substitutions are those which are conservative in nature, i.e., those substitutions that take place within a family of amino acids that are related in their side chains. Specifically, amino acids are generally divided into four families: (1) acidic: aspartate and glutamate; (2) basic: lysine, arginine, and histidine; (3) non-polar: alanine, valine, leucine, isoleucine, proline, phenylalanine, methionine, tryptophan; and (4) uncharged polar: glycine, asparagine, glutamine,

cystine, serine, theronine, tyrosine. Phenylalanine, tryptophan, and tyrosine are sometimes classified as aromatic amino acids. For example, it is reasonably predictable that an isolated replacement of leucine with isoleucine or valine, an asparate with a glutamate, a threonine with a serine, or a similar conservative replacement of an amino acid with a structurally related amino acid, will not have a major effect on the biological activity. Proteins having substantially the same amino acid sequence as the reference molecule, but possessing minor amino acid substitutions that do not substantially affect the immunogenicity of the protein, are therefore within the definition of the reference polypeptide.

The VLPs of the invention can be formed from any viral protein, particle-forming polypeptide derived from the viral protein, or combination of viral proteins or fragments thereof, that have the capability of forming particles under appropriate conditions. The requirements for the particle-forming viral proteins are that if the particle is formed in the cytoplasm of the host cell, the protein must be sufficiently stable in the host cell in which it is expressed such that formation of virus-like structures will result, and that the polypeptide will automatically assemble into a virus-like structure in the cell of the recombinant expression system used. If the protein is secreted into culture media, conditions can be adjusted such that VLPs will form. Furthermore, the particle-forming protein should not be cytotoxic in the expression host and should not be able to replicate in the host in which the VLP will be used.

Preferred particle-forming polypeptides include coronavirus M and E proteins, preferably SARS M and E proteins.

Methods and suitable conditions for forming particles from a wide variety of viral proteins are known in the art. VLPs have been produced, for example from proteins derived from influenza virus (such as HA or NA), Hepatitis B virus (such as core or capsid proteins), Hepatitis E virus, measles virus, Sindbis virus, Rotavirus, Foot-and-Mouth Disease virus, Retrovirus, Norwalk virus, human Papilloma virus, HIV, RNA-phages, Q8-phage (such as coat proteins), GA-phage, fr-phage, AP205 phage, and Ty (such as retrotransposon Ty protein p1). VLPs are discussed further in WO 03/024480, WO 03/024481, and Niikura et al., Virology (2002) 293:273-280; Lenz et al., J. Immunology (2001) 5246-5355; Pinto, et al., J. Infectious Diseases (2003) 188:327-338; and Gerber et al., J. Virology (2001) 75(10):4752-4760.

As explained above, VLPs can spontaneously form when the particle-forming polypeptide of interest is recombinantly expressed in an appropriate host cell. Thus, the VLPs for use in the present invention may be prepared using recombinant techniques, well known in the art. In this regard, genes encoding the particle-forming polypeptide in question can be isolated from DNA libraries or directly from cells and tissues containing the same, using known techniques. The genes encoding the particle-forming polypeptides can also be produced synthetically, based on the known sequences. The nucleotide sequence can be designed with the appropriate codons for

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the particular amino sequence desired. In general, one will select preferred codons for the intended host in which the sequence will be expressed (e.g. human codons for human DNA vaccines). The complete sequence is generally assembled from overlapping oligonucleotides prepared by standard methods and assembled into a complete coding sequence. See., e.g., Edge, Nature (1981) 292:756; Nambair et al. Science (1984) 223:1299; Jay et al., J. Biol. Chem. (1984) 259:6311.

Once the coding sequences for the desired particle-forming polypeptides have been isolated or synthesized, they can be cloned into any suitable vector or replicon for expression. Numerous cloning vectors are known to those of skill in the art, and the selection of an appropriate cloning vector is a matter of choice. See, generally, Sambrook et al. The vector is then used to transform an appropriate host cell. Suitable expression systems include, but are not limited to, bacterial, mammalian, bacuolvirus/insect, vaccinia, Semliki Forest virus (SFV), yeast, and Xenopus expression systems, well known in the art.

A number of cell lines suitable for use as host cells for producing the VLPs of the invention are known in the art. Suitable mammalian cell lines include, but are not limited to, Chinese Hamster Ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (COS), human hepatocellular carcinoma cells (e.g., Hep G2), Madin-Darby bovine kidney ("MDBK") cells, as well as others. Mammalian sources of cells include, but are not limited to, human or non-human primate (e.g., MRC-5 (ATCC CCL-171), WI-38 (ATCC CCL-75), HUH, human embryonic kidney cells (293 cells, typically transformed by sheared adenovirus type 5 DNA), VERO cells from monkey kidneys (including, for example COS7 cells), horse, cow (e.g., MDBK cells), sheep, dog (e.g., MDCK cells from dog kidneys, ATCC CCL34 MDCK (NBL2) or MDCK 33016, deposit number DSM ACC 2219 as described in WO 97/37001), cat, and rodent (e.g., hamster cells such as BHK21-F, HKCC cells, or Chinese hamster ovary cells (CHO cells)), and may be obtained from a wide variety of developmental stages, including for example, adult, neonatal, fetal, and embryo.

Bacterial hosts suitable for production of VLPs of the invention include E. coli, Bacillus subtilis, and Streptoccocus spp. Yeast hosts suitable for production of VLPs of the invention include Saccharomyces cerevisiae, Candida albicans, Candida maltosa, Hansenula polymorpha, Kluyveromyces fragilis, Kluyveromyces lactis, Pichia guillerimondii, Pichia pastoris, Schizosaccharomyces pombe and Yarrowia lipolytica. Insect cells suitable for production of VLPs of the invention (i.e., via baculovirus expression vectors) include Aedes aegypti, Autographa californica, Bombyx mori, Drosophila melanogaster, Spodptera frugiperda, and Trichoplusia ni.

Viral vectors can be used for the production of particles in eukaryotic cells, such as those derived from the pox family of viruses, including vaccinia virus and avian poxvirus. Additional,

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vaccinia based infection/transfection systems, such as those as described in Tomei et al., J. Virol (1993) 67:4017-4026 and Selby et al., J. Gen. Virol. (1993) 74:1103-1113, can also be used to generate the VLPs of the invention. In this system, cells are first transfected in vitro with a vaccinia virus recombinant that encodes the bacteriophage T7 RNA polymerase. This polymerase only transcribes templates bearing T7 promoters. Following infection, cells are transfected with the DNA of interest, driven by a T7 promoter. The polymerase expressed in the cytoplasm from the vaccinia virus recombinant transcribes the transfected DNA into RNA which is then translated into protein by the host translation machinery. The method provides for high level, transient, cytoplasmic production of large quantities of RNA and its translation products.

Depending on the expression system and host selected, the VLPs are produced by growing host cells transformed by an expression vector under conditions whereby the particle-forming polypeptide is expressed and VLPs can be formed. The selection of the appropriate growth conditions is within the skill of the art. If the VLPs are formed intracellularly, the cells are then disrupted, using chemical, physical or mechanical means, which lyse the cells yet keep the VLPs substantially intact. Such methods are known the those of skill in the art and are described in, e.g., Protein Purification Applications: A Practical Approach, (E.L.V. Harris and S. Angal, Eds., 1990).

The particles are then isolated using methods that preserve the integrity thereof, such as by gradient centrifugation, e.g., cesium chloride (CsCl) and sucrose gradients, and the like (see, e.g., Kirnbauer et al., J. Virol. (1993) 67:6929-6936), ion exchange chromatography (including anion exchange chromatography such as DMAE and TMAE), hydroxyapatitic chromatography (see WO 00/09671), hydrophobic interaction chromatography, gel filtration chromatography and other filtration methods such as nanometric filtration and ultrafiltration. Preferably at least one anion exchange step is performed during purification, and more preferably at least two anion exchange steps are used.

VLP formulations of the invention may be further processed by methods known in the art to disassemble the VLPs into smaller, protein containing moieties using a high concentration of reducing agent, followed by reassembly of the VLPs by either removal of the reducing agent or by addition of excess oxidant. The resulting reassembled VLPs may have improved homogeneity, stability and immunogenic properties. In addition, further therapeutic or prophylactic agents may be formulated into the VLPs upon reassembly. See McCarthy et al., J. Virology (1998) 72(1):32-41. See also WO 99/13056 and WO 01/42780. Reducing agents suitable for use in VLP disassembly include sulfhydryl reducing agents (such as glutathion, beta mercaptoethanol, dithiothreitol, dithioerythritol, cysteine, hydrogen sulfide and mixtures thereof) preferably contained in moderate to low ionic strength buffers. Sufficient exposure time of the VLPs to the reducing agent will be required to achieve a suitable amount of VLP disassembly.

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Adjuvants may be added to the VLPs of the invention to enhance the immunogenicity of the SARS viral antigens. Antigens suitable for use with VLPs include those described, supra. For example, the VLPs of the invention may be adsorbed onto an aluminum adjuvant

The VLPs of the invention may formulated to enhance their stability. Additional components which may enhance the stability of a VLP formulation include salts, buffers, non-ionic surfactants and other stabilizers such as polymeric polyanion stabilizers. See WO 00/45841.

The ionic strength of a solution comprising VLP particles may be maintained by the presence of salts. Almost any salt which can contribute to the control of the ionic strength may be used. Preferred salts which can be used to adjust ionic strength include physiologically acceptable salts such as NaCl, KCl, Na₂SO₄, (NH₄)₂SO₄, sodium phosphate and sodium citrate. Preferably, the salt component is present in concentrations of from about 0.10 M to 1 M. Very high concentrations are not preferred due to the practical limitations of parenteral injection of high salt concentrations. Instead, more moderate salt concentrations, such as more physiological concentrations of about 0.15M to about 0.5M with 0.15M-0.32M NaCl are preferred.

Buffers may also be used to enhance the stability of the VLP formulations of the invention. Preferably, the buffer optimizes the VLP stability while maintaining the pH range so that the vaccine formulation will not be irritating to the recipient. Buffers preferably maintain the pH of the vaccine formulation within a range of p/H 5.5-7.0, more preferably 6.0-6.5. Buffers suitable for vaccine formulations are known in the art and include, for example, histidine and imidazole. Preferably, the concentration of the buffer will range from about 2mM to about 100 mM, more preferably 5 mM to about 20 mM. Phosphate containing buffers are generally not preferred when the VLP is adsorbed or otherwise formulated with an aluminum compound.

Non-ionic surfactants may be used to enchance the stability of the VLP formulations of the invention. Surfactants suitable for use in vaccine formulations are known in the art and include, for example, polyoxyethylene sorbital fatty acid esters (Polysorbates) such as Polysorbate 80 (e.g., TWEEN 80), Polysorbate 20 (e.g., TWEEN 20), polyoxyethylene alkyl ethers (e.g., Brij 35, Brij 58), as well as others, including Triton X-100, Triton X-114, NP-40, Span 85 and the Pluronic series of non-ionic surfactants (e.g., Pluronic 121). The surfactant is preferably present in a concentration of from about 0.0005% to about 0.5% (wt/vol).

Polymeric polyanion stabilizers may also be used to enchance the stability of the VLP formulations of the invention. Suitable polymeric polyanionic stabilizers for use in the invention comprise either a single long chain or multiple cross linked chains; either type possessing multiple negative charges along the chains when in solution. Examples of suitable polyanionic polymers include proteins, polyanions, peptides and polynucelic acids. Specific examples include carboxymethyl cellulose, heparin, polyamino acids (such as poly(Glu), poly(Asp), and

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Poly (Glu, Phe), oxidized glutathione, polynuceltodies, RNA, DNA and serum albumins. The concentration of the polmeric polyanion stabilizers is preferably from about 0.01% to about 0.5%, particularly about 0.05-0.1% (by weight).

G. Passive Immunization via Antibodies to the SARS Antigens of the Invention

The invention includes antibodies specific to the SARS antigens of the invention and methods of treatment or prevention of SARS virus related disease by administrating an effective amount of SARS antibodies to a mammalian subject. Antibodies specific the SARS antigens can be produced by one skilled in the art. Preferably, the antibodies are specific to the spike (S) protein of the SARS virus. Potent neutralization of the SARS coronavirus using a human monoclonal anti-spike antibody has been reported (Sui et al. (2004) PNAS USA 101:2536-2541). A IgG1 form of the monoclonal antibody showed a higher affinity (1.59 nM) than a scFv form (32.3 nM).

The antibodies of the invention are specific and selective to SARS antigens.

In one embodiment, the antibodies of the invention are generated by administering a SARS antigen to an animal. The method may also include isolating the antibodies from the animal.

The antibodies of the invention may be polyclonal or monoclonal antibody preparations, monospecific antisera, human antibodies, or may be hybrid or chimeric antibodies, such as humanized antibodies, altered antibodies (Fab')₂ fragments, F(ab) fragments, Fv fragments, single-domain antibodies, dimeric or trimeric antibody fragments or constructs, minibodies, or functional fragments thereof which bind to the antigen in question.

Antibodies are produced using techniques well known to those of skill in the art and disclosed in, for example, US Patent Nos. 4,011,308; 4,722,890; 4,016,043; 3,876,504; 3,770,380; and 4,372,745. For example, polyclonal antibodies are generated by immunizing a suitable animal, such as a mouse, rat, rabbit, sheep, or goat, with an antigen of interest. In order to enhance immunogenicity, the antigen can be linked to a carrier prior to immunization. Such carriers are well known to those of ordinary skill in the art. Immunization is generally performed by mixing or emulsifying the antigen in saline, preferably in an adjuvant such as Freund's complete adjuvant, and injecting the mixture or emulsion parenterally (generally subcutaneously or intramuscularly). The animal is generally boosted 2-6 weeks later with one or more injections of the antigen in saline, preferably using Freund's incomplete adjuvant. Antibodies may also be generated by in vitro immunization, using methods known in the art. Polyclonal antiserum is then obtained from the immunized animal.

Monoclonal antibodies are generally prepared using the method of Kohler & Milstein (1975) Nature 256:495-497, or a modification thereof. Typically, a mouse or rat is immunized as described above. Rabbits may also be used. However, rather than bleeding the animal to extract serum, the spleen (and optionally several large lymph nodes) is removed and dissociated into

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single cells. If desired, the spleen cells may be screened (after removal of non-specifically adherent cells) by applying a cell suspension to a plate or well coated with the antigen. B-cells, expressing membrane-bound immunoglobulin specific for the antigen, will bind to the plate, and are not rinsed away with the rest of the suspension. Resulting B-cells, or all dissociated spleen cells, are then induced to fuse with myeloma cells to form hybridomas, and are cultured in a selective medium (e.g., hypoxanthine, aminopterin, thymidine medium, "HAT"). The resulting hybridomas are plated by limiting dilution, and are assayed for the production of antibodies which bind specifically to the immunizing antigen (and which do not bind to unrelated antigens). The selected monoclonal antibody-secreting hybridomas are then cultured either in vitro (e.g., in tissue culture bottles or hollow fiber reactors), or in vivo (e.g., as ascites in mice).

Humanized and chimeric antibodies are also useful in the invention. Hybrid (chimeric) antibody molecules are generally discussed in Winter et al. (1991) Nature 349: 293-299 and US Patent No. 4,816,567. Humanized antibody molecules are generally discussed in Riechmann et al. (1988) Nature 332:323-327; Verhoeyan et al. (1988) Science 239:1534-1536; and U.K. Patent Publication No. GB 2,276,169, published 21 September 1994). One approach to engineering a humanized antibody involves cloning recombinant DNA containing the promoter, leader, and variable-region sequences from a mouse antibody gene and the constant-region exons from a human antibody gene to create a mouse-human chimera, a humanized antibody. See generally, Kuby, "Immunology, 3rd Edition", W.H. Freeman and Company, New York (1998) at page 136.

Antibody fragments which retain the ability to recognize a SARS antigen are also included within the scope of the invention. A number of antibody fragments are known in the art which comprise antigen-binding sites capable of exhibiting immunological binding properties of an intact antibody molecule. For example, functional antibody fragments can be produced by cleaving a constant region, not responsible for antigen binding, from the antibody molecule, using e.g., pepsin, to produce F(ab')₂ fragments. These fragments will contain two antigen binding sites, but lack a portion of the constant region from each of the heavy chains. Similarly, if desired, Fab fragments, comprising a single antigen binding site, can be produced, e.g., by digestion of polyclonal or monclonal antibodies with papain. Functional fragments, including only the variable regions of the heavy and light chains, can also be produced, using standard techniques such as recombinant production or preferential proteolytic cleavage of immunoglobulin molecules. These fragments are known as F_v. See, e.g., Inbar et al. (1972) Proc. Nat. Acad. Sci USA 69:2659-2662; Hochman et al. (1976) Biochem 15:2706-2710; and Ehrlich et al. (1980) Biochem 19:4091-4096.

A single-chain Fv ("sFv" or scFv") polypeptide is a covalently linked $V_{H^-}V_L$ heterodimer which is expressed from a gene fusion including V_{H^-} and V_{L^-} encoding genes linked by a peptide-

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encoding linker. Huston et al. (1988) Proc. Nat. Acad. Sci. USA 85:5879-5883. A number of methods have been described to discern and develop chemical strucutres (linkers) for converting the naturally aggregated, but chemically separated, light and heavy polypeptide chains from an antibody V region into an sFv molecule which will fold into a three dimensional structure substantially similar to the structure of an antigen-binding site. See, e.g., US Patent Nos. 5,091,513; 5,132,405; and 4,946,778. The sFv molecules may be produced using methods described in thea rt. See, e.g., Huston et al. (1988) Proc. Nat. Acad. Sci USA 85:5879-5338; US Patent Nos. 5,091,513; 5,132,405 and 4,946,778. Design criteria include determining the appropriate length to span the distance between the C-terminus of one chain and the N-terminus of the other, wherein the linker is generally formed from small hydrophilic amino acid residues that do not coil or form secondary structures. Such methods have been described in the art. See, e.g., US Patent Nos. 5,091,513; 5,132,405 and 4,946,778. Suitable linkers generally comprise polypeptide chains of alternating sets of glycine and serine residues, and may include glutamic acid and lysine residues inserted to enhance solubility. Anti-spike scFv antibodies have been reported (Sui et al. (2004) PNAS USA 101:2536-2541).

"Mini-antibodies" or "minibodies" will also find use with the present invention.

Minibodies are sFv polypeptide chains which include oligomerization domains at their Ctermini, separated from the sFv by a hinge region. Pack et al., (1992) Biochem 31:1579-1584.

The oligomerization domain comprises self-associating α-helices, e.g., leucine zippers, that can
be further stabilized by additional disulfide bonds. The oligomerization domain is designed to be
compatible with vectorial folding across a membrane, a process thought to facilitate in vivo
folding of the polypeptide into a functional binding protein. Generally, minibodies are produced
using recombinant methods well known in the art. See, e.g., Pack et al., (1992) Biochem
31:1579-1584; Cumber et al. (1992) J. Immunology 149B:120-126.

Non-conventional means can also be used to generate and identify the antibodies of the invention. For example, a phage display library can be screened for antibodies which bind to the SARS antigens of the invention. See generally, Siegel, "Recombinant Monoclonal Antibody Technology", Transfus. Clin. Biol. (2002) 2(1): 15-22; Sidhu, "Phage Display in Pharmaceutical Biotechnology", Curr. Opin. Biotechnol. (2000) 11(6):610-616; Sharon, et.al., "Recombinant Polyclonal Antibody Libraries", Comb. Chem. High Throughput Screen (2000) 3(3): 185-196; and Schmitz et al., "Phage Display: A Molecular Tool for the Generation of Antibodies-Review", Placenta, (2000) 21 SupplA: S106-12.

The antibodies of the invention may also be generated by administering the polynucleotide sequence encoding for the SARS antigen into an animal. The SARS antigen is then expressed in vivo, and antibodies specific to the SARS antigen are generated *in vivo*. Methods for polynucleotide delivery of the SARS antigens of the invention are discussed in section 4 below.

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The antibodies of the invention are preferably specific to the SARS virus.

H. Combinations of one or more of any of the above approaches in a vaccine

The compositions of the invention further comprise combinations of one or more of the compositions discussed above. For instance, the invention comprises a composition comprising an attenuated SARS virus and a subunit SARS viral antigen.

I. Combinations of SARS antigens and other Respiratory Virus Antigens

The invention further relates to vaccine formulations comprising one or more SARS virus antigens and one or more other respiratory virus antigens. Additional respiratory virus antigens suitable for use in the invention include antigens from influenza virus, human rhinovirus (HRV), parainfluenza virus (PIV), respiratory syncytial virus (RSV), adenovirus, metapneumovirus, and rhinovirus. The additional respiratory virus antigen could also be from a coronavirus other than the SARS coronavirus, such as the NL63 human coronavirus (van der Hoek et al. (2004) Nature Medicine 10:368-373). Preferably, the additional respiratory virus antigen is an influenza viral antigen.

The invention may also comprise one or more bacterial or viral antigens in combination with the SARS viral antigen. Antigens may be used alone or in any combination. (See, e.g., WO 02/00249 describing the use of combinations of bacterial antigens). The combinations may include multiple antigens from the same pathogen, multiple antigens from different pathogens or multiple antigens from the same and from different pathogens. Thus, bacterial, viral, and/or other antigens may be included in the same composition or may be administered to the same subject separately. It is generally preferred that combinations of antigens be used to raise an immune response be used in combinations.

Non-limiting examples of bacterial pathogens which may be used in the invention include diphtheria (See, e.g., Chapter 3 of Vaccines, 1998, eds. Plotkin & Mortimer (ISBN 0-7216-1946-0), staphylococcus (e.g., Staphylococcus aureus as described in Kuroda et al. (2001) Lancet 357:1225-1240), cholera, tuberculosis, C. tetani, also known as tetanus (See, e.g., Chapter 4 of Vaccines, 1998, eds. Plotkin & Mortimer (ISBN 0-7216-1946-0), Group A and Group B streptococcus (including Streptococcus pneumoniae, Streptococcus agalactiae and Streptococcus pyogenes as described, for example, in Watson et al. (2000) Pediatr. Infect. Dis. J. 19:331-332; Rubin et al. (2000) Pediatr Clin. North Am. 47:269-284; Jedrzejas et al. (2001) Microbiol Mol Biol Rev 65:187-207; Schuchat (1999) Lancet 353:51-56; GB patent applications 0026333.5; 0028727.6; 015640.7; Dale et al. (1999) Infect Dis Clin North Am 13:227-1243; Ferretti et al. (2001) PNAS USA 98:4658-4663), pertussis (See, e.g., Gusttafsson et al. (1996) N. Engl. J. Med. 334:349-355; Rappuoli et al. (1991) TIBTECH 9:232-238), meningitis, Moraxella catarrhalis (See, e.g., McMichael (2000) Vaccine 19 Suppl. 1:S101-107) and other pathogenic states,

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including, without limitation, Neisseria meningitides (A, B, C, Y), Neisseria gonorrhoeae (See, e.g., WO 99/24578; WO 99/36544; and WO 99/57280), Helicobacter pylori (e.g., CagA, VacA, NAP, HopX, HopY and/or urease as described, for example, WO 93/18150; WO 99/53310; WO 98/04702) and Haemophilus influenza. Hemophilus influenza type B (HIB) (See, e.g.,

5 Costantino et al. (1999) Vaccine 17:1251-1263), Porphyromonas gingivalis (Ross et al. (2001) Vaccine 19:4135-4132) and combinations thereof.

Non-limiting examples of viral pathogens which may be used in the invention include meningitis, rhinovirus, influenza (Kawaoka *et al.*, *Virology* (1990) 179:759-767; Webster *et al.*, "Antigenic variation among type A influenza viruses," p. 127-168. In: P. Palese and D.W.

- Kingsbury (ed.), Genetics of influenza viruses. Springer-Verlag, New York), respiratory syncytial virus (RSV), parainfluenza virus (PIV), rotavirus (e.g., VP1, VP2, VP3, VP4, VP6, VP7, NSP1, NSP2, NSP3, NSP4 or NSP5 and other rotavirus antigens, for example as described in WO 00/26380) and the like. Antigens derived from other viruses will also find use in the present invention, such as without limitation, proteins from members of the families
- Picomaviridae (e.g., polioviruses, etc. as described, for example, in Sutter et al. (2000) Pediatr Clin North Am 47:287-308; Zimmerman & Spann (1999) Am Fam Physician 59:113-118; 125-126); Caliciviridae; Togaviridae (e.g., rubella virus, etc.); Flaviviridae, including the genera flavivirus (e.g., yellow fever virus, Japanese encephalitis virus, serotypes of Dengue virus, tick borne encephalitis virus, West Nile virus, St. Louis encephalitis virus); pestivirus (e.g., classical porcine fever virus, bovine viral diarrhea virus, border disease virus); and hepacivirus (e.g.,
 - hepatitis A, B and C as described, for example, in US Patent Nos. 4,702,909; 5,011,915; 5,698,390; 6,027,729; and 6,297,048); Parvovirus (e.g., parvovirus B19); Coronaviridae; Reoviridae; Rimaviridae; Rhabodoviridae (e.g., rabies virus, etc. as described for example in Dressen et al. (1997) Vaccine 15 Suppl:s2-6; MMWR Morb Mortal Wkly Rep. 1998 Jan 16:47(1):12, 19); Filoviridae; Paramyxoviridae (e.g., numps virus, measles virus, respiratory
 - syncytial virus, etc. as described in Chapters 9 to 11 of Vaccines, 1998, eds. Plotkin & Mortimer (ISBN 0-7216-1946-0); Orthomyxoviridae (e.g., influenza virus types A, B and C, etc. as described in Chapter 19 of Vaccines, 1998, eds. Plotkin & Mortimer (ISBN 0-7216-1946-0),); Bunyaviridae; Arenaviridae; Retroviradae (e.g., HTLV-1; HTLV-11; HIV-1 (also known as
- HTLV-III, LAV, ARV, HTI,R, etc.)), including but not limited to antigens from the isolates HIVIIIb, HIVSF2, HIVLAV, HIVI-AL, I-IIVMN, SF162); HIV-I CM235, HIV-I US4; HIV-2; simian immunodeficiency virus (SIV) among others. Additionally, antigens may also be derived from human papilloma virus (HPV) and the tick-borne encephalitis viruses. See, e.g. Virology, 3rd Edition (W.K. Joklik ed. 1988); Fundamental Virology, 2nd Edition (B.N. Fields and D.M.
- Knipe, eds, 1991), for a description of these and other viruses.

Proteins may also be derived from the herpesvirus family, including proteins derived from herpes simplex virus (HSV) types 1 and 2, such as HSV-1 and HSV-2 glycoproteins gB, gD and gH; antigens derived from varicella zoster virus (VZV), Epstein-Barr virus (EBV) and cytomegalovirus (CMV) including CMV gB and gH (See, US Patent No. 4,689,225 and PCT Publication WO 89/07143): and antigens derived from other human herpesviruses such as HHV6 and HHV7. (See, e.g. Chee et al., Cytomegaloviruses (J.K. McDougall, ed., Springer-Verlag 1990) pp. 125-169, for a review of the protein coding content of cytomegalovirus; McGeoch et al., J. Gen. Virol. (1988) 69:1531-1574, for a discussion of the various HSV-1 encoded proteins; US Patent No. 5,171,568 for a discussion of HSV-1 and HSV-2 gB and gD proteins and the genes encoding therefor; Baer et al., Nature (1984) 310:207-211, for the identification of protein coding sequences in an EBV genome; and Davison and Scott, J. Gen. Virol. (1986) 67:1759-1816, for a review of VZV). Herpes simplex virus (HSV) rgD2 is a recombinant protein produced in genetically engineered Chinese hamster ovary cells. This protein has the normal anchor region truncated, resulting in a glycosylated protein secreted into tissue culture medium. The gD2 can be purified in the CHO medium to greater than 90% purity. Human immunodeficiency virus (HIV) env-2-3 is a recombinant form of the HIV enveloped protein produced in genetically engineered Saccharomyces cerevisae. This protein represents the entire protein region of HIV gp120 but is non-glycosylated and denatured as purified from the yeast. HIV gpl20 is a fully glycosylated, secreted form of gp120 produced in CHO cells in a fashion similar to the gD2 above. Additional HSV antigens suitable for use in immunogenic compositions are described in PCT Publications W0 85/04587 and W0 88/02634, the disclosures of which are incorporated herein by reference in their entirety. Mixtures of gB and gD antigens, which are truncated surface antigens lacking the anchor regions, are particularly preferred.

Antigens from the hepatitis family of viruses, including hepatitis A virus (HAV) (See, e.g., Bell et al. (2000) Pediatr Infect Dis. J. 19:1187-1188; Iwarson (1995) APMIS 103:321-326), hepatitis B virus (HBV) (See, e.g., Gerlich et al. (1990) Vaccine 8 Suppl:S63-68 & 79-80), hepatitis C virus (HCV) (See, e.g., PCT/US88/04125, published European application number 318216), the delta hepatitis virus (HDV), hepatitis E virus (HEV) and hepatitis G virus (HGV), can also be conveniently used in the techniques described herein. By way of example, the viral genomic sequence of HCV is known, as are methods for obtaining the sequence. See, e.g., International Publication Nos. WO 89/04669; WO 90/11089; and WO 90/14436. Also included in the invention are molecular variants of such polypeptides, for example as described in PCT/US99/31245; PCT/US99/31273 and PCT/US99/31272. The HCV genome encodes several viral proteins, including E1 (also known as E) and E2 (also known as E2/NSI) and an N-terminal nucleocapsid protein (termed "core") (see, Houghton et al., Hepatology (1991) 14:381-388, for a discussion of HCV proteins, including E1 and E2). Similarly, the sequence for the δ-antigen

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from HDV is known (see, e.g., US Patent No. 5,378,814) and this antigen can also be conveniently used in the present composition and methods. Additionally, antigens derived from HBV, such as the core antigen, the surface antigen, SAg, as well as the presurface sequences, pre-S1 and pre-S2 (formerly called pre-S), as well as combinations of the above, such as SAg/pre-S1, SAg/pre-S2, SAg/pre-S1/pre-S2, and pre-S1/pre-S2, will find use herein. See, e.g., "HBV Vaccines - from the laboratory to license: a case study" in Mackett, M. and Williamson, J.D., Human Vaccines and Vaccination, pp. 159-176, for a discussion of HBV structure; and US Patent Nos. 4,722,840, 5,098,704, 5,324,513, incorporated herein by reference in their entireties; Beames et al., J. Virol. (1995) 69:6833-6838, Birnbaum et al., J. Virol. (1990) 64:3319-3330; and Zhou et al., J. Virol. (1991) 65:5457-5464. Each of these proteins, as well as antigenic fragments thereof, will find use in the present composition and methods.

Influenza virus is another example of a virus for which the present invention will be particularly useful. Specifically, the envelope glycoproteins HA and NA of influenza A are of particular interest for generating an immune response. Numerous HA subtypes of influenza A have been identified (Kawaoka et al., Virology (1990) 179:759-767; Webster et al., "Antigenic variation among type A influenza viruses," p. 127-168. In: P. Palese and D.W. Kingsbury (ed.), Genetics of influenza viruses. Springer-Verlag, New York). Thus, proteins derived from any of these isolates can also be used in the compositions and methods described herein.

Non-limiting examples of parasitic antigens include those derived from organisms causing malaria and Lyme disease.

The methods of the invention comprise administering an immunogenic composition comprising a SARS viral antigen (including one or more of an inactivated SARS virus, an attenuated SARS virus, a split SARS virus preparation or a recombinant or purified subunit formulation of one or more SARS viral antigens) to an animal. The immunogenic compositions used in the invention can comprise an immunologically effective amount of the SARS viral antigen. An "immunologically effective amount" is an amount sufficient to allow the mammal to raise an immune response to the SARS antigen.

The immune response preferably involves the production of antibodies specific to the SARS antigen. The amount of antibodies produced will vary depending on several factors including the animal used, the presence of an adjuvant, etc.

The immunogenic compositions of the invention may further comprise one or more adjuvants.

The immunogenic compositions of the invention may be administered mucosally. Suitable routes of mucosal administration include oral, intranasal, intragastric, pulmonary, intestinal, rectal, ocular and vaginal routes. The immunogenic composition may be adapted for mucosal administration. For instance, where the composition is for oral administration, it may be in the

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form of tablets or capsules, optionally enteric-coated, liquid, transgenic plants, etc. Where the composition is for intranasal administration, it may be in the form of a nasal spray, nasal drops, gel or powder.

The immunogenic compositions of the invention may be administered parenterally. Suitable routes of parenteral administration include intramuscular (IM), subcutaneous, intravenous, intraverional, intradermal, transcutaneous, and transdermal (see e.g., International patent application WO 98/20734) routes, as well as delivery to the interstitial space of a tissue. The immunogenic composition may be adapted for parenteral administration, for instance in the form of an injectable that may be sterile and pyrogen free.

Vaccines of the invention may be administered in conjunction with other immunoregulatory agents. In particular, compositions will usually include an adjuvant. Preferred further adjuvants include, but are not limited to, one or more of the following set forth below:

A. Mineral Containing Compositions

Mineral containing compositions suitable for use as adjuvants in the invention include mineral salts, such as aluminium salts and calcium salts. The invention includes mineral salts such as hydroxides (e.g. oxyhydroxides), phosphates (e.g. hydroxyphoshpates, orthophosphates), sulphates, etc. (e.g. see chapters 8 & 9 of Vaccine design: the subunit and adjuvant approach (1995) Powell & Newman. ISBN 0-306-44867-X.), or mixtures of different mineral compounds, with the compounds taking any suitable form (e.g. gel, crystalline, amorphous, etc.), and with adsorption being preferred. The mineral containing compositions may also be formulated as a particle of metal salt. See WO00/23105.

B. Oil-Emulsions

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Oil-emulsion compositions suitable for use as adjuvants in the invention include squalene-water emulsions, such as MF59 (5% Squalene, 0.5% Tween 80, and 0.5% Span 85, formulated into submicron particles using a microfluidizer). See WO90/14837. See also, Frey et al., "Comparison of the safety, tolerability, and immunogenicity of a MF59-adjuvanted influenza vaccine and a non-adjuvanted influenza vaccine in non-elderly adults", Vaccine (2003) 21:4234-4237.

Particularly preferred adjuvants for use in the compositions are submicron oil-inwater emulsions. Preferred submicron oil-in-water emulsions for use herein are squalene/water emulsions optionally containing varying amounts of MTP-PE, such as a submicron oil-in-water emulsion containing 4-5% w/v squalene, 0.25-1.0% w/v Tween 80 TM (polyoxyelthylenesorbitan monooleate), and/or 0.25-1.0% Span 85TM (sorbitan trioleate), and, optionally, N-acetylmuramyl-L-alanyl-D-isogluatminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-

huydroxyphosphophoryloxy)-ethylamine (MTP-PE), for example, the submicron oil-in-water

emulsion known as "MF59" (International Publication No. WO 90/14837; US Patent Nos 6.299.884 and 6.451.325, incorporated herein by reference in their entireties: and Ott et al. "MF59 -- Design and Evaluation of a Safe and Potent Adjuvant for Human Vaccines" in Vaccine Design: The Subunit and Adjuvant Approach (Powell, M.F. and Newman, M.J. eds.) Plenum Press, New York, 1995, pp. 277-296). MF59 contains 4-5% w/v Squalene (e.g., 4.3%), 0.25-5 0.5% w/v Tween 80™, and 0.5% w/v Span 85™ and optionally contains various amounts of MTP-PE, formulated into submicron particles using a microfluidizer such as Model 110Y microfluidizer (Microfluidics, Newton, MA). For example, MTP-PE may be present in an amount of about 0-500 μg/dose, more preferably 0-250 μg/dose and most preferably, 0-100 lO µg/dose. As used herein, the term "MF59-0" refers to the above submicron oil-in-water emulsion lacking MTP-PE, while the term MF59-MTP denotes a formulation that contains MTP-PE. For instance, "MF59-100" contains 100 µg MTP-PE per dose, and so on. MF69, another submicron oil-in-water emulsion for use herein, contains 4.3% w/v squalene, 0.25% w/v Tween 80™, and 0.75% w/v Span 85™ and optionally MTP-PE. Yet another submicron oil-in-water emulsion is MF75, also known as SAF, containing 10% squalene, 0.4% Tween 80™, 5% 5 pluronic-blocked polymer L121, and thr-MDP, also microfluidized into a submicron emulsion. MF75-MTP denotes an MF75 formulation that includes MTP, such as from 100-400 µg MTP-PE per dose.

Submicron oil-in-water emulsions, methods of making the same and immunostimulating agents, such as muramyl peptides, for use in the compositions, are described in detail in International Publication No. WO 90114837 and US Patent Nos. 6,299,884 and 6,45 1,325, incorporated herein by reference in their entireties.

Complete Freund's adjuvant (CFA) and incomplete Freund's adjuvant (IFA) may also be used as adjuvants in the invention.

C. Saponin Formulations

Saponin formulations, may also be used as adjuvants in the invention. Saponins are a heterologous group of sterol glycosides and triterpenoid glycosides that are found in the bark, leaves, stems, roots and even flowers of a wide range of plant species. Saponin from the bark of the Quillaia saponaria Molina tree have been widely studied as adjuvants. Saponin can also be commercially obtained from Smilax ornata (sarsaprilla), Gypsophilla paniculata (brides veil), and Saponaria officianalis (soap root). Saponin adjuvant formulations include purified formulations, such as QS21, as well as lipid formulations, such as ISCOMs.

Saponin compositions have been purified using High Performance Thin Layer

Chromatography (HP-LC) and Reversed Phase High Performance Liquid Chromatography (RP-HPLC). Specific purified fractions using these techniques have been identified, including QS7,

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QS17, QS18, QS21, QH-A, QH-B and QH-C. Preferably, the saponin is QS21. A method of production of QS21 is disclosed in US Patent No. 5,057,540. Saponin formulations may also comprise a sterol, such as cholesterol (see WO 96/33739).

Combinations of saponins and cholesterols can be used to form unique particles called Immunostimulating Complexs (ISCOMs). ISCOMs typically also include a phospholipid such as phosphatidylethanolamine or phosphatidylcholine. Any known saponin can be used in ISCOMs. Preferably, the ISCOM includes one or more of Quil A, QHA and QHC. ISCOMs are further described in EP 0 109 942, WO 96/11711 and WO 96/33739. Optionally, the ISCOMS may be devoid of additional detergent. See WO00/07621.

A review of the development of saponin based adjuvants can be found at Barr, et al., "ISCOMs and other saponin based adjuvants", Advanced Drug Delivery Reviews (1998) 32:247-271. See also Sjolander, et al., "Uptake and adjuvant activity of orally delivered saponin and ISCOM vaccines", Advanced Drug Delivery Reviews (1998) 32:321-338.

D. Bacterial or Microbial Derivatives

Adjuvants suitable for use in the invention include bacterial or microbial derivatives such as:

(1) Non-toxic derivatives of enterobacterial lipopolysaccharide (LPS)

Such derivatives include Monophosphoryl lipid A (MPL) and 3-O-deacylated MPL (3dMPL). 3dMPL is a mixture of 3 De-O-acylated monophosphoryl lipid A with 4, 5 or 6 acylated chains. A preferred "small particle" form of 3 De-O-acylated monophosphoryl lipid A is disclosed in EP 0 689 454. Such "small particles" of 3dMPL are small enough to be sterile filtered through a 0.22 micron membrane (see EP 0 689 454). Other non-toxic LPS derivatives include monophosphoryl lipid A mimics, such as aminoalkyl glucosaminide phosphate derivatives e.g. RC-529. See Johnson et al. (1999) Bioorg Med Chem Lett 9:2273-2278.

(2) Lipid A Derivatives

Lipid A derivatives include derivatives of lipid A from Escherichia coli such as OM-174. OM-174 is described for example in Meraldi et al., "OM-174, a New Adjuvant with a Potential for Human Use, Induces a Protective Response with Administered with the Synthetic C-Terminal Fragment 242-310 from the circumsporozoite protein of Plasmodium berghei", Vaccine (2003) 21:2485-2491; and Pajak, et al., "The Adjuvant OM-174 induces both the migration and maturation of murine dendritic cells in vivo", Vaccine (2003) 21:836-842.

(3) Immunostimulatory oligonucleotides

Immunostimulatory oligonucleotides suitable for use as adjuvants in the invention include nucleotide sequences containing a CpG motif (a sequence containing an unmethylated cytosine followed by guanosine and linked by a phosphate bond). Bacterial double stranded RNA or

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oligonucleotides containing palindromic or poly(dG) sequences have also been shown to be immunostimulatory.

The CpG's can include nucleotide modifications/analogs such as phosphorothioate modifications and can be double-stranded or single-stranded. Optionally, the guanosine may be replaced with an analog such as 2'-deoxy-7-deazaguanosine. See Kandimalla, et al., "Divergent synthetic nucleotide motif recognition pattern: design and development of potent immunomodulatory oligodeoxyribonucleotide agents with distinct cytokine induction profiles", Nucleic Acids Research (2003) 31(9): 2393-2400; WO 02/26757 and WO 99/62923 for examples of possible analog substitutions. The adjuvant effect of CpG oligonucleotides is further discussed in Krieg, "CpG motifs: the active ingredient in bacterial extracts?", Nature Medicine (2003) 9(7): 831-835; McCluskie, et al., "Parenteral and mucosal prime-boost immunization strategies in mice with hepatitis B surface antigen and CpG DNA", FEMS Immunology and Medical Microbiology (2002) 32:179-185; WO 98/40100; US Patent No. 6,207,646; US Patent No. 6,239,116 and US Patent No. 6,429,199.

The CpG sequence may be directed to TLR9, such as the motif GTCGTT or TTCGTT. See Kandimalla, et al., "Toll-like receptor 9: modulation of recognition and cytokine induction by novel synthetic CpG DNAs", Biochemical Society Transactions (2003) 31 (part 3): 654-658. The CpG sequence may be specific for inducing a Th1 immune response, such as a CpG-A ODN, or it may be more specific for inducing a B cell response, such a CpG-B ODN. CpG-A and CpG-B ODNs are discussed in Blackwell, et al., "CpG-A-Induced Monocyte IFN-gamma-Inducible Protein-10 Production is Regulated by Plasmacytoid Dendritic Cell Derived IFN-alpha", J. Immunol. (2003) 170(8):4061-4068; Krieg, "From A to Z on CpG", TRENDS in Immunology (2002) 23(2): 64-65 and WO 01/95935. Preferably, the CpG is a CpG-A ODN.

Preferably, the CpG oligonucleotide is constructed so that the 5' end is accessible for receptor recognition. Optionally, two CpG oligonucleotide sequences may be attached at their 3' ends to form "immunomers". See, for example, Kandimalla, et al., "Secondary structures in CpG oligonucleotides affect immunostimulatory activity", BBRC (2003) 306:948-953; Kandimalla, et al., "Toll-like receptor 9: modulation of recognition and cytokine induction by novel synthetic GpG DNAs", Biochemical Society Transactions (2003) 31(part 3):664-658; Bhagat et al., "CpG penta- and hexadeoxyribonucleotides as potent immunomodulatory agents" BBRC (2003) 300:853-861 and WO 03/035836.

(4) ADP-ribosylating toxins and detoxified derivatives thereof.

Bacterial ADP-ribosylating toxins and detoxified derivatives thereof, may be used as adjuvants in the invention. Preferably, the protein is derived from *E. coli* (i.e., E. coli heat labile enterotoxin "LT), cholera ("CT"), or pertussis ("PT"). The use of detoxified ADP-ribosylating toxins as mucosal adjuvants is described in WO 95/17211 and as parenteral adjuvants in WO

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98/42375. Preferably, the adjuvant is a detoxified LT mutant such as LT-K63, LT-R72, and LTR192G. The use of ADP-ribosylating toxins and detoxified derivaties thereof, particularly LT-K63 and LT-R72, as adjuvants can be found in the following references, each of which is specifically incorporated by reference herein in their entirety: Beignon, et al., "The LTR72 Mutant of Heat-Labile Enterotoxin of Escherichia coli Enahnces the Ability of Peptide Antigens to Elicit CD4+ T Cells and Secrete Gamma Interferon after Coapplication onto Bare Skin", Infection and Immunity (2002) 70(6):3012-3019; Pizza, et al., "Mucosal vaccines: non-toxic derivatives of LT and CT as mucosal adjuvants", Vaccine (2001) 19:2534-2541; Pizza, et al., "LTK63 and LTR72, two mucosal adjuvants ready for clinical trials" Int. J. Med. Microbiol (2000) 290(4-5):455-461; Scharton-Kersten et al., "Transcutaneous Immunization with Bacterial ADP-Ribosylating Exotoxins, Subunits and Unrelated Adjuvants", Infection and Immunity (2000) 68(9):5306-5313; Ryan et al., "Mutants of Escherichia coli Heat-Labile Toxin Act as Effective Mucosal Adjuvants for Nasal Delivery of an Acellular Pertussis Vaccine: Differential Effects of the Nontoxic AB Complex and Enzyme Activity on Th1 and Th2 Cells" Infection and Immunity (1999) 67(12):6270-6280; Partidos et al., "Heat-labile enterotoxin of Escherichia coli and its site-directed mutant LTK63 enhance the proliferative and cytotoxic T-cell responses to intranasally co-immunized synthetic peptides", Immunol. Lett. (1999) 67(3):209-216; Peppoloni et al., "Mutants of the Escherichia coli heat-labile enterotoxin as safe and strong adjuvants for intranasal delivery of vaccines", Vaccines (2003) 2(2):285-293; and Pine et al., (2002) "Intranasal immunization with influenza vaccine and a detoxified mutant of heat labile enterotoxin from Escherichia coli (LTK63)" J. Control Release (2002) 85(1-3):263-270. Numerical reference for amino acid substitutions is preferably based on the alignments of the A and B subunits of ADP-ribosylating toxins set forth in Domenighini et al., Mol. Microbiol (1995) 15(6):1165-1167, specifically incorporated herein by reference in its entirety.

E. Human Immunomodulators

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Human immunomodulators suitable for use as adjuvants in the invention include cytokines, such as interleukins (e.g. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12, etc.), interferons (e.g. interferon-γ), macrophage colony stimulating factor, and tumor necrosis factor.

F. Bioadhesives and Mucoadhesives

Bioadhesives and mucoadhesives may also be used as adjuvants in the invention. Suitable bioadhesives include esterified hyaluronic acid microspheres (Singh et al. (2001) J. Cont. Rele. 70:267-276) or mucoadhesives such as cross-linked derivatives of poly(acrylic acid), polyvinyl alcohol, polyvinyl pyrollidone, polysaccharides and carboxymethylcellulose. Chitosan and derivatives thereof may also be used as adjuvants in the invention. E.g., WO99/27960.

G. Microparticles

Microparticles may also be used as adjuvants in the invention. Microparticles (i.e. a particle of ~100nm to ~150 μ m in diameter, more preferably ~200nm to ~30 μ m in diameter, and most preferably ~500nm to ~10 μ m in diameter) formed from materials that are biodegradable and non-toxic (e.g. a poly(α -hydroxy acid), a polyhydroxybutyric acid, a polyorthoester, a polyanhydride, a polycaprolactone, etc.), with poly(lactide-co-glycolide) are preferred, optionally treated to have a negatively-charged surface (e.g. with SDS) or a positively-charged surface (e.g. with a cationic detergent, such as CTAB).

H. Liposomes

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Examples of liposome formulations suitable for use as adjuvants are described in US Patent No. 6,090,406, US Patent No. 5,916,588, and EP 0 626 169.

I. Polyoxyethylene ether and Polyoxyethylene Ester Formulations

Adjuvants suitable for use in the invention include polyoxyethylene ethers and polyoxyethylene esters. WO99/52549. Such formulations further include polyoxyethylene sorbitan ester surfactants in combination with an octoxynol (WO01/21207) as well as polyoxyethylene alkyl ethers or ester surfactants in combination with at least one additional nonionic surfactant such as an octoxynol (WO01/21152).

Preferred polyoxyethylene ethers are selected from the following group: polyoxyethylene-9-lauryl ether (laureth 9), polyoxyethylene-9-steoryl ether, polyoxyethylene-8-steoryl ether, polyoxyethylene-4-lauryl ether, polyoxyethylene-35-lauryl ether, and polyoxyethylene-23-lauryl ether.

J. Polyphosphazene (PCPP)

PCPP formulations are described, for example, in Andrianov et al., "Preparation of hydrogel microspheres by coacervation of aqueous polyphophazene solutions", Biomaterials (1998) 19(1-3):109-115 and Payne et al., "Protein Release from Polyphosphazene Matrices", Adv. Drug. Delivery Review (1998) 31(3):185-196.

K. Muramyl peptides

Examples of muramyl peptides suitable for use as adjuvants in the invention include N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetyl-normuramyl-L-alanyl-D-isoglutamine (nor-MDP), and N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphosphoryloxy)-ethylamine MTP-PE).

L. Imidazoquinolone Compounds.

Examples of imidazoquinolone compounds suitable for use adjuvants in the invention include Imiquamod and its homologues, described further in Stanley, "Imiquimod and the

imidazoquinolones: mechanism of action and therapeutic potential" Clin Exp Dermatol (2002) 2<u>T</u>(7):571-577 and Jones, "Resiquimod 3M", Curr Opin Investig Drugs (2003) <u>4</u>(2):214-218.

M. Virosomes and Virus Like Particles (VLPs)

Virosomes and Virus Like Particles (VLPs) can also be used as adjuvants in the invention. These structures generally contain one or more proteins from a virus optionally combined or formulated with a phospholipid. They are generally non-pathogenic, non-replicating and generally do not contain any of the native viral genome. The viral proteins may be recombinantly produced or isolated from whole viruses. These viral proteins suitable for use in virosomes or VLPs include proteins derived from influenza virus (such as HA or NA). Hepatitis B virus (such as core or capsid proteins), Hepatitis E virus, measles virus, Sindbis virus, Rotavirus, Foot-and-Mouth Disease virus, Retrovirus, Norwalk virus, human Papilloma virus, HIV, RNA-phages, OB-phage (such as coat proteins), GA-phage, fr-phage, AP205 phage, and Ty (such as retrotransposon Ty protein p1). VLPs are discussed further in WO 03/024480, WO 03/024481. and Niikura et al., "Chimeric Recombinant Hepatitis E Virus-Like Particles as an Oral Vaccine Vehicle Presenting Foreign Epitopes", Virology (2002) 293:273-280; Lenz et al., "Papillomarivurs-Like Particles Induce Acute Activation of Dendritic Cells", Journal of Immunology (2001) 5246-5355; Pinto, et al., "Cellular Immune Responses to Human Papillomavirus (HPV)-16 L1 Healthy Volunteers Immunized with Recombinant HPV-16 L1 Virus-Like Particles", Journal of Infectious Diseases (2003) 188:327-338; and Gerber et al., "Human Papillomavrisu Virus-Like Particles Are Efficient Oral Immunogens when Coadministered with Escherichia coli Heat-Labile Entertoxin Mutant R192G or CpG", Journal of Virology (2001) 75(10):4752-4760. Virosomes are discussed further in, for example, Gluck et al., "New Technology Platforms in the Development of Vaccines for the Future", Vaccine (2002) 20:B10-B16.

The invention may also comprise combinations of aspects of one or more of the adjuvants identified above. For example, the following adjuvant compositions may be used in the invention:

- (1) a saponin and an oil-in-water emulsion (WO99/11241);
- (2) a saponin (e.g.., QS21) + a non-toxic LPS derivative (e.g., 3dMPL) (see WO 94/00153);
 - (3) a saponin (e.g., QS21) + a non-toxic LPS derivative (e.g., 3dMPL) + a cholesterol;
 - (4) a saponin (e.g. QS21) + 3dMPL + IL-12 (optionally + a sterol) (WO98/57659);
 - (5) combinations of 3dMPL with, for example, QS21 and/or oil-in-water emulsions (See European patent applications 0835318, 0735898 and 0761231);

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(6) SAF, containing 10% Squalane, 0.4% Tween 80, 5% pluronic-block polymer L121, and thr-MDP, either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion.

- (7) Ribi™ adjuvant system (RAS), (Ribi Immunochem) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (Detox™); and
- (8) one or more mineral salts (such as an aluminum salt) + a non-toxic derivative of LPS (such as 3dPML).

Aluminium salts and MF59 are preferred adjuvants for parenteral immunisation. Mutant bacterial toxins are preferred mucosal adjuvants.

As mentioned above, adjuvants suitable for use in the invention may also include one or more of the following:

- E.coli heat-labile enterotoxin ("LT"), or detoxified mutants thereof, such as the K63 or R72 mutants;
 - cholera toxin ("CT"), or detoxified mutants thereof;
 - microparticles (i.e., a particle of ~100nm to ~150 μ m in diameter, more preferably ~200nm to ~30 μ m in diameter, and most preferably ~500nm to ~10 μ m in diameter) formed from materials that are biodegradable and non-toxic (e.g. a poly(α -hydroxy acid), a polyhydroxybutyric acid, a polyorthoester, a polyanhydride, a polycaprolactone etc.):
 - a polyoxyethylene ether or a polyoxyethylene ester (see International patent application WO 99/52549);
- a polyoxyethylene sorbitan ester surfactant in combination with an octoxynol (see International patent application WO 01/21207) or a polyoxyethylene alkyl ether or ester surfactant in combination with at least one additional non-ionic surfactant such as an octoxynol (see International patent application WO 01/21152);
 - chitosan (e.g. International patent application WO 99/27960)
- an immunostimulatory oligonucleotide (e.g. a CpG oligonucleotide) and a saponin (see International patent application WO 00/62800)
 - immunostimulatory double stranded RNA.
- aluminum compounds (e.g. aluminum hydroxide, aluminum phosphate, aluminum hydroxyphosphate, oxyhydroxide, orthophosphate, sulfate etc. (e.g. see chapters 8 & 9 of Vaccine design: the subunit and adjuvant aproach, eds. Powell & Newman, Plenum Press 1995 (ISBN 0-306-44867-X) (hereinafter "Vaccine design"), or mixtures of different aluminum compounds, with the compounds taking any suitable form (e.g. gel, crystalline, amorphous etc.), and with adsorption being preferred;

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- MF59 (5% Squalene, 0.5% Tween 80, and 0.5% Span 85, formulated into submicron particles using a microfluidizer) (see Chapter 10 of Vaccine design; see also International patent application WO 90/14837);

- liposomes (see Chapters 13 and 14 of Vaccine design);
- ISCOMs (see Chapter 23 of Vaccine design);
- SAF, containing 10% Squalane, 0.4% Tween 80, 5% plurohic-block polymer L121, and thr-MDP, either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion (see Chapter 12 of *Vaccine design*);
- Ribi™ adjuvant system (RAS), (Ribi Immunochem) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (Detox™);
- saponin adjuvants, such as QuilA or QS21 (see Chapter 22 of Vaccine design), also known as StimulonTM;
 - ISCOMs, which may be devoid of additional detergent (WO 00/07621):
 - complete Freund's adjuvant (CFA) and incomplete Freund's adjuvant (IFA);
- cytokines, such as interleukins (e.g. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12, etc.), interferons (e.g. interferon-γ), macrophage colony stimulating factor, tumor necrosis factor, etc. (see Chapters 27 & 28 of *Vaccine design*);
- monophosphoryl lipid A (MPL) or 3-O-deacylated MPL (3dMPL) (e.g. chapter 21 of $\it Vaccine\ design$);
- combinations of 3dMPL with, for example, QS21 and/or oil-in-water emulsions (European patent applications 0835318, 0735898 and 0761231);
- oligonucleotides comprising CpG motifs (see Krieg (2000) Vaccine, 19:618-622; Krieg (2001) Curr. Opin. Mol. Ther., 2001, 3:15-24; WO 96/02555, WO 98/16247, WO 98/18810, WO 98/40100, WO 98/55495, WO 98/37919 and WO 98/52581, etc.) i.e. containing at least one CG dinucleotide,
- a polyoxyethylene ether or a polyoxyethylene ester (International patent application WO99/52549);
- a polyoxyethylene sorbitan ester surfactant in combination with an octoxynol (International patent application WO 01/21207) or a polyoxyethylene alkyl ether or ester surfactant in combination with at least one additional non-ionic surfactant such as an octoxynol (WO 01/21152);
- an immunostimulatory oligonucleotide (e.g. a CpG oligonucleotide) and a saponin (WO00/62800);

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- an immunostimulant and a particle of metal salt (International patent application WO00/23105);

- a saponin and an oil-in-water emulsion (WO 99/11241);
- a saponin (e.g. QS21) + 3dMPL + IL-12 (optionally + a sterol) (WO 98/57659).

Other adjuvants suitable for mucosal or parenteral administration are also available (e.g. see chapter 7 of Vaccine design: the subunit and adjuvant aproach, eds. Powell & Newman, Plenum Press 1995 (ISBN 0-306-44867-X).

Mutants of LT are preferred mucosal adjuvants, in particular the "K63" and "R72" mutants (e.g. see International patent application WO 98/18928), as these result in an enhanced immune response.

Microparticles are also preferred mucosal adjuvants. These are preferably derived from a poly(a-hydroxy acid), in particular, from a poly(lactide) ("PLA"), a copolymer of D,L-lactide and glycolide or glycolic acid, such as a poly(D,L-lactide-co-glycolide) ("PLG" or "PLGA"), or a copolymer of D,L-lactide and caprolactone. The microparticles may be derived from any of various polymeric starting materials which have a variety of molecular weights and, in the case of the copolymers such as PLG, a variety of lactide:glycolide ratios, the selection of which will be largely a matter of choice, depending in part on the coadministered antigen.

The SARS virus (inactivated or attenuated), viral antigens, antibodies or adjuvants of the invention may be entrapped within the microparticles, or may be adsorbed to them. Entrapment within PLG microparticles is preferred. PLG microparticles are discussed in further detail in Morris et al., (1994), Vaccine, 12:5-11, in chapter 13 of Mucosal Vaccines, eds. Kiyono et al., Academic Press 1996 (ISBN 012410587), and in chapters 16 & 18 of Vaccine design: the subunit and adjuvant aproach, eds. Powell & Newman, Plenum Press 1995 (ISBN 0-306-44867-X).

LT mutants may advantageously be used in combination with microparticle-entrapped antigen, resulting in significantly enhanced immune responses.

Aluminium compounds and MF59 are preferred adjuvants for parenteral use.

The composition may include an antibiotic.

The immunogenic compositions of the invention may be administered in a single dose, or as part of an administration regime. The regime may include priming and boosting doses, which may be administered mucosally, parenterally, or various combinations thereof.

The methods of the invention further comprise treating or preventing a SARS virus-related disease by administering to an animal a composition comprising an effective amount of the antibodies of the invention. An "effective amount" of the antibodies of the invention is an amount sufficient to provide passive immunization protection or treatment to the animal. Preferably, the antibodies of the invention are specific to the SARS viral antigen.

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Methods of treatment may combine both immunogenic compositions and antibody compositions. Accordingly the invention comprises a method for treating or preventing a SARS virus-related disease comprising administering an immunogenic composition comprising an immunologically effective amount of a SARS viral antigen and administering an effective amount of antibodies specific to SARS viral antigen. The immunogenic composition and the antibodies may be administered together or separately. The invention further comprises a composition comprising an immunologically effective amount of a SARS viral antigen and further comprising an effective amount of antibodies specific to a SARS viral antigen.

The SARS viral antigens and antibodies of the invention may also be administered in polynucleotide form. The SARS viral antigens and/or antibody proteins are then expressed in vivo.

The SARS viral antigens and the antibodies of the invention can also be delivered using one or more gene vectors, administered via nucleic acid immunization or the like using standard gene delivery protocols. Methods for gene delivery are known in the art. Seé, e.g., US Patent Nos. 5,399,346, 5,580,859, 5,589,466. The constructs can be delivered (e.g., injected) either subcutaneously, epidermally, intradermally, intramuscularly, intravenous, mucosally (such as nasally, rectally and vaginally), intraperitoneally, orally or combinations thereof. Intramuscular injection of 25µg plasmid DNA encoding spike antigens, in 200µl PBS pH 7.4, at weeks 0, 3 and 6, has been described for mice by Yang et al. (2004) Nature 428:561-564.

An exemplary replication-deficient gene delivery vehicle that may be used in the practice of the present invention is any of the alphavirus vectors, described in, for example, US Patent Nos. 6,342,372; 6,329,201 and International Publication WO 01/92552.

A number of viral based systems have been developed for gene transfer into mammalian cells. For example, retroviruses provide a convenient platform for gene delivery systems. Selected sequences can be inserted into a vector and packaged in retroviral particles using techniques known in the art. The recombinant virus can then be isolated and delivered to cells of the subject either in vivo or ex vivo. A number of retroviral systems have been described (US Patent No. 5,219,740; Miller & Rosman, BioTechniques (1989) 7:980-990; Miller, A.D., Human Gene Therapy (1990) 1:5-14; Scarpa et al., Virology (1991) 180:849-852; Burns et al., Proc. Natl. Acad. Sci. USA (1993) 90:8033-8037; and Boris-Lawrie & Temin, Cur. Opin. Genet. Develop. (1993) 3:102-109.

A number of adenovirus vectors have also been described. Unlike retroviruses which integrate into the host genome, adenoviruses persist extrachromosomally thus minimizing the risks associated with insertional mutagenesis (Haj-Ahmad and Graham, *J. Virol.* (1986) 57:267-274; Bett et al., *J. Virol.* (1993) 67:5911-5921; Mittereder et al., Human Gene Therapy (1994)

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5:717-729; Seth et al., J. Virol. (1994) 68:933-940; Barr et al., Gene Therapy (1994) 1:51-58; Berkner, K.L. BioTechniques (1988) 6:616-629; and Rich et al., Human Gene Therapy (1993) 4:461-476). Adenoviral delivery of codon-optimsed versions of the genes encoding SARS coronavirus structural antigens spike S1, membrane protein and nucleocapsid protein has been investigated in rhesus macaques and found to invoke a strong neutralizing antibody response (Gao et al. (2003) Lancet 362(9399):1895-1896).

Additionally, various adeno-associated virus (AAV) vector systems have been developed for gene delivery. AAV vectors can be readily constructed using techniques well known in the art. See, e.g., US Patent Nos. 5,173,414 and 5,139,941; International Publication Nos. WO 92/01070 (published 23 January 1992) and WO 93/03769 (published 4 March 1993); Lebkowski et al., Molec. Cell. Biol. (1988) 8:3988-3996; Vincent et al., Vaccines 90 (1990) (Cold Spring Harbor Laboratory Press); Carter, B.J. Current Opinion in Biotechnology (1992) 3:533-539; Muzyczka, N. Current Topics in Microbiol. and Immunol. (1992) 158:97-129; Kotin, R.M. Human Gene Therapy (1994) 5:793-801; Shelling and Smith, Gene Therapy (1994) 1:165-169; and Zhou et al., J. Exp. Med. (1994) 179:1867-1875.

Another vector system useful for delivering polynucleotides, mucosally and otherwise, is the enterically administered recombinant poxvirus vaccines described by Small, Jr., P.A., et al. (US Patent No. 5,676,950, issued October 14, 1997, herein incorporated by reference) as well as the vaccinia virus and avian poxviruses. By way of example, vaccinia virus recombinants expressing the genes can be constructed as follows. The DNA encoding the SARS antigen or antibody or antibody coding sequence is first inserted into an appropriate vector so that it is adjacent to a vaccinia promoter and flanking vaccinia DNA sequences, such as the sequence encoding thymidine kinase (TK). This vector is then used to transfect cells that are simultaneously infected with vaccinia. Homologous recombination serves to insert the vaccinia promoter plus the gene encoding the coding sequences of interest into the viral genome. The resulting TK recombinant can be selected by culturing the cells in the presence of 5-bromodeoxyuridine and picking viral plaques resistant thereto.

Alternatively, avipoxviruses, such as the fowlpox and canarypox viruses, can also be used to deliver genes encoding the SARS viral antigens or antibodies of the invention. Recombinant avipox viruses, expressing immunogens from mammalian pathogens, are known to confer protective immunity when administered to non-avian species. The use of an avipox vector is particularly desirable in human and other mammalian species since members of the avipox genus can only productively replicate in susceptible avian species and therefore are not infective in mammalian cells. Methods for producing recombinant avipoxviruses are known in the art and employ genetic recombination, as described above with respect to the production of vaccinia

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viruses. See, e.g., WO 91/12882; WO 89/03429; and WO 92/03545. Picornavirus-derived vectors can also be used. (See, e.g., US Patent Nos. 5,614,413 and 6,063,384).

Molecular conjugate vectors, such as the adenovirus chimeric vectors described in Michael et al., J. Biol. Chem. (1993) 268:6866-6869 and Wagner et al., Proc. Natl. Acad. Sci. USA (1992) 89:6099-6103, can also be used for gene delivery.

A vaccinia based infection/transfection system can be conveniently used to provide for inducible, transient expression of the coding sequences of interest (for example, a SARS viral antigen or antibody expression cassette) in a host cell. In this system, cells are first infected in vitro with a vaccinia virus recombinant that encodes the bacteriophage T7 RNA polymerase. This polymerase displays exquisite specificity in that it only transcribes templates bearing T7 promoters. Following infection, cells are transfected with the polynucleotide of interest, driven by a T7 promoter. The polymerase expressed in the cytoplasm from the vaccinia virus recombinant transcribes the transfected DNA into RNA that is then translated into protein by the host translational machinery. The method provides for high level, transient, cytoplasmic production of large quantities of RNA and its translation products. See, e.g., Elroy-Stein and Moss, Proc. Natl. Acad. Sci. USA (1990) 87:6743-6747; Fuerst et al., Proc. Natl. Acad. Sci. USA (1986) 83:8122-8126.

As an alternative approach to infection with vaccinia or avipox virus recombinants, or to the delivery of genes using other viral vectors, an amplification system can be used that will lead to high level expression following introduction into host cells. Specifically, a T7 RNA polymerase promoter preceding the coding region for T7 RNA polymerase can be engineered. Translation of RNA derived from this template will generate T7 RNA polymerase that in turn will transcribe more template. Concomitantly, there will be a cDNA whose expression is under the control of the T7 promoter. Thus, some of the T7 RNA polymerase generated from translation of the amplification template RNA will lead to transcription of the desired gene. Because some T7 RNA polymerase is required to initiate the amplification, T7 RNA polymerase can be introduced into cells along with the template(s) to prime the transcription reaction. The polymerase can be introduced as a protein or on a plasmid encoding the RNA polymerase. For a further discussion of T7 systems and their use for transforming cells, see, e.g., International Publication No. WO 94/26911; Studier and Moffatt, J. Mol. Biol. (1986) 189:113-130; Deng and Wolff, Gene (1994) 143:245-249; Gao et al., Biochem. Biophys. Res. Commun. (1994) 200:1201-1206; Gao and Huang, Nuc. Acids Res. (1993) 21:2867-2872; Chen et al., Nuc. Acids Res. (1994) 22:2114-2120; and US Patent No. 5,135,855.

The immunogenic compositions of the invention may further comprise diluents, such as water, saline, glycerol, ethanol, etc. Additionally, auxiliary substances, such as wetting or

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emulsifying agents, pH buffering substances, and the like may be included in the immunogenic composition.

The immunogenic compositions used in the invention can be administered to an animal. Animals suitable for use in the methods of the invention include humans and other primates, including non-human primates such as chimpanzees, and other apes and monkey species; farm animals such as cattle, sheep, pigs, goats and horses, domestic animals such as dogs and cats; laboratory animals including rodents such as mice, rats and guinea pigs; birds, including domestic, wild and game birds such as chickens, turkeys and other gallinaceous birds, ducks, geese and the like. Animals suitable for use in the invention can be of any age, including both adult and newborn. Transgenic animals can also be used in the invention.

The immunogenic compositions of the invention can be used to treat or prevent SARS virus-related diseases.

The compositions of the invention are preferably pharmaceutically acceptable and pharmacologically acceptable. In particularly, the compositions are preferably not biologically or otherwise undesirable, i.e., the material may be administered to an individual in a formulation or composition without causing any undesirable biological effects or interacting in a deleterious manner with any of the components of the composition in which it is contained.

Pharmaceutically acceptable salts can also be used in compositions of the invention, for example, mineral salts such as hydrochlorides, hydrobromides, phosphates, or sulfates, as well as salts of organic acids such as acetates, proprionates, malonates, or benzoates. Especially useful protein substrates are serum albumins, keyhole limpet hemocyanin, immunoglobulin molecules, thyroglobulin, ovalbumin, tetanus toxoid, and other proteins well known to those of skill in the art. Compositions of the invention can also contain liquids or excipients, such as water, saline, glycerol, dextrose, ethanol, or the like, singly or in combination, as well as substances such as wetting agents, emulsifying agents, or pH buffering agents. Liposomes can also be used as a carrier for a composition of the invention.

SARS specific reagents and analytical assays may be used in the manufacture and testing of the vaccines of the invention. Such analytical assays include, for example: 1) virus titration and plaque assays for quantitation of infectious virus particles, 2) a neutralization assay with constant virus and varying serum dilutions, 3) a two step RT-PCR system (Light Cycler-Roche) for detection of negative strand viral RNA, with the target sequence located within the N gene, providing highest possible sensitivity, and 4) ELISA and western blot assays for detection and qualification of viral proteins.

In addition, rabbit polyclonal antiserum has been generated to obtain antibody reagents (and demonstrate induction of neutralizing antibodies) against the SARS-CoV. A sample protocol for generating such reagents is set forth below. The virus is first cultivated in suitable

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cell culture, such as Vero cells, and pelleted through a 20% sucrose (w/v) cushion. The pellet is then subjected to a glycerol potassium-tartrate gradient for further purification. The virus-containing fraction is then diluted and pelleted by ultracentrifugation. The pellet is then dissolved in PBS and the virus is inactivated with C₃H₄O₂ (beta-propiolactone, BPL). Two rabbits are immunized subcutaneously (SC) on day 0, 14, and 28 with 1x10⁹ inactivated viral particles mixed with IFA as adjuvant. Rabbits are bled on days 0 (pre-inoculation), 13, 28, and 35 (I week after 3rd immunization). Sera obtained from this protocol were tested for their reactivity against SARS-CoV proteins in western blots and found to react with the major structural proteins spike (S), membrane (M), and nucleocapsid (N).

J. Emerging coronavirus vaccines

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The SARS epidemic has lead to increased awareness of viral infections caused by coronaviruses. The vaccines of the invention may be adapted to prevent or treat emerging strains of coronavirus, including emerging strains of SARS virus.

The invention provides a vaccine comprising an inactivated (or killed) human coronavirus, an attenuated human coronavirus, a split human coronavirus preparation, or a recombinant or purified subunit formulation of one or more antigens from a human coronavirus, wherein the human coronavirus is not the SARS coronavirus. Optionally, the human coronavirus is not the 229E coronavirus. Optionally, the human coronavirus is not the NL63 coronavirus. Thus the invention provides a vaccine as defined above, wherein the human coronavirus is not the SARS coronavirus, is not the 229E coronavirus, is not the OC43 coronavirus and is not the NL63 coronavirus. Such vaccines are useful for preventing and/or treating emerging human coronavirus infections.

The invention also provides a vaccine comprising: (a) an inactivated (or killed) human coronavirus, an attenuated human coronavirus, a split human coronavirus preparation, or a recombinant or purified subunit formulation of one or more antigens from a human coronavirus, wherein the human coronavirus is not the SARS coronavirus, as defined above; and (b) an inactivated (or killed) human coronavirus, an attenuated human coronavirus, a split human coronavirus preparation, or a recombinant or purified subunit formulation of one or more antigens from a human coronavirus, wherein the human coronavirus is the SARS coronavirus. Such vaccines are useful for preventing and/or treating both SARS and other human coronaviruses.

As well as providing vaccines comprising antigens from more than one type of coronavirus, the invention also provides vaccines comprising antigens from more than one strain of the same coronavirus *e.g.* different strains of the SARS coronavirus, or different strains of a coronavirus other than the SARS coronavirus. In one embodiment, the vaccine comprises antigens from at least two strains of coronavirus, or at least three strains of coronavirus. In one

embodiment, the vaccine comprises antigens from at least two types of coronavirus. In one embodiment, the vaccine comprises at least one antigen from each of the known types of coronaviruses (type I, type II and type III). Such vaccines follow the model of current influenza vaccines.

The selection of coronaviruses and/or coronavirus strains for use in vaccines of the invention can be based on various criteria. For instance, selection may be based on viruses and/or strains that have been detected in the geographical region (e.g. northern or southern hemisphere, a particular country, etc.) where the vaccine targeted. Selection may be based on the results of animal surveillance e.g. of viruses detected in cat populations. Selection may be based on the results of clinical surveillance e.g. of viruses detected in patients hospitalized with respiratory infection. Selection may be performed every year e.g. prior to winter. Vaccines may also be administered yearly, again following the model of current influenza vaccines.

Preferred vaccines are sufficiently immunogenic to provide a neutralizing immune response, and more preferably a protective and/or therapeutic immune response. Particularly preferred vaccines meet the efficacy requirements that may be specified by the WHO from time to time.

A preferred subunit antigen for inclusion in vaccines of the invention is a purified spike protein, more preferably in oligomeric (e.g. trimeric) form. The spike protein may or my not be cleaved e.g. into its S1 and S2 products.

The techniques disclosed above for selecting viruses and/or strains for production of vaccines can also be used to select appropriate viruses and/or strains from which HR1 and HR2 sequences can be obtained for providing therapeutic peptides, as disclosed above.

III. DIAGNOSTIC COMPOSITIONS AND METHODS OF THE INVENTION

The invention provides methods for detecting the SARS coronavirus. Detection in patient samples can be used to detect and diagnose infections by the virus. Detection in donated blood can be used to prevent inadvertent transmission of the virus during blood transplant procedures Detection methods fall into three main categories: detection of SARS virus nucleic acids; detection of SARS virus proteins; and detection of anti-SARS virus immune responses. The invention provides all such methods.

As used herein when referring to nucleotide sequences, particularly oligonucleotide probes and primers, "similar" sequences includes those sequences that are at least 90% identical to known SARSV genomic sequence and includes sequences that are at least 95% identical, at least 99% identical and 100% identical to the SARSV genomic sequence over the length of the probe or primer.

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To a slurry of LiAlH₄ (4 equivalents) in dry THF, cooled to 0 °C under N_2 , was slowly added drop-wise a solution of methyl 4-[4-(1-BOC-piperazin-4-yl)phenoxymethyl]benzoate (1 equivalent) in dry THF. Once the addition was complete, the slurry was heated to reflux at 80 °C for 1 hour. The slurry was subsequently cooled to 0 °C and treated with water, 10% aq. NaOH and with water again. The resulting solids were filtered, and the filtrate was diluted with chloroform, washed with brine, dried over MgSO₄ and concentrated, providing the crude 4-[4-(4-methylpiperazin-1-yl)phenoxymethyl]benzyl alcohol that was used without purification.

To a solution of DMSO (2.6 equivalents) in dry DCM, cooled to -78 °C under N_2 was added oxalyl chloride (1.1 equivalents) in DCM drop-wise. The solution was stirred at -78 °C for 5 minutes before a solution of 4-[4-(4-methylpiperazin-1-yl)phenoxymethyl]benzyl alcohol (1 equivalent) in DCM was added drop-wise, and allowed to stir at -78 °C for another 30 minutes. Triethylamine (2.5 equivalents) was slowly dripped in before allowing the solution to reach ambient temperatures. The solution was washed with aqueous NaHCO₃ and brine, dried over MgSO₄ and concentrated to provide the crude 4-[4-(4-methylpiperazin-1-yl)phenoxymethyl]benzaldehyde that was converted to thiosemicarbazones according to Scheme 7.

PYRROLES

Scheme 15

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Synthesis of Pyrrole

10 Preparation of tert-butyl (2E)-3-(2,4-dichlorophenyl)prop-2-enoate (2).

Neat DIC (1.4 eq) was added to a well stirred solution of cinnamate (1 eq), t-butyl alcohol (4 eq), DMAP (1.4 eq) and CH_2Cl_2 under argon at rt. (Note - The cinnamate must be completely in solution that may require gentle warming. Allow the solution to cool to room temperature before adding the DIC. To avoid an exotherm on larger scales, it may be prudent to -270-

dilute the DIC with CH_2Cl_2 before the addition and have an ice bath ready.) After stirring for 8 hours, the reaction develops a white precipitate. The reaction may be monitored by TLC eluting with 25% EtOAc/Hexane (R_f of product was 0.9). The entire reaction was loaded into a separatory funnel (washing with CH_2Cl_2). The organic mixture was washed with citrate, sat. aq. NaHCO₃, water, and brine. The organic layer was dried (Na_2SO_4), filtered, and concentrated to dryness to give the crude product as an oil. The crude oil was mixed with hexane and stirred for 30 min. The precipitate that forms was filtered over celite and the filtrate was evaporated. The hexane mixture was loaded onto a filter plug of silica and eluted with EtOAc/hexane (97:2 v/v). The first eluted UV active fractions are collected and evaporated to give >99% pure 2 (75-80% yields).

Preparation of tert-butyl 4-(2,4-dichlorophenyl)pyrrole-3-carboxylate (3).

Dry ether was added to NaH (1.5 eq as the oil dispersion) under argon. After decanting off the ether via syringe, the NaH was suspended again with fresh ether under argon. A solution of TOSMIC (1.1 eq) and 2 (1 eq) dissolved in a mixture of ether and DMSO was added dropwise to the stirred suspension of NaH at 0 °C over 20-30 min. The addition was mildly exothermic and evolved gas. After the addition, the reaction was allowed to warm to ambient rt. The progress of the reaction was followed by TLC (25% EtOAc/Hexane, the UV active product was at $R_f = 0.4$) and LCMS until done (-2-3 h). Upon completion, the reaction was carefully quenched with sat. aq. NH4cI (added slowly to avoid strong gas evolution and exotherm) and diluted with ether. The layers were separated and the organic phase was washed with sat. aq. NaHCO₃, water, and brine. The crude dark solid can be purified by recrystallization. Best results were achieved either through recrystallization directly from a mixture of hot EtOAc/hexane (1:3 ν) or by dissolving the crude product in minimal hot EtOAc followed by addition of hexane (-2 volumes of hexane based on the volume of EtOAc). The hot solutions were allowed to cool to room temperature and age over night. The crystals were first filtered and then washed with hexane giving 99% pure product in 60-70 % yield.

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Preparation of *tert*-butyl 4-(2,4-dichlorophenyl)-1-[3-(1,3-dioxobenzo[c]azolin-2-yl)propyl]pyrrole-3-carboxylate (4).

Solid NaH (1.5 eq as the oil dispersion) was added in small portions to a solution of pyrrole 3 (1 eq) and 3-bromopropyl phthalimide (1.2 eq) dissolved in DMF stirred at room temperature and flushed with argon. NOTE - Some gas evolves, but the temperature does not seem to rise above 40-50 °C. The reaction was stirred for 1.5 h at room temperature under argon. The reaction was followed by TLC (CH₂Cl₂/acetonitrile (95:5 v/v), the UV active product was at R_f = 0.5) and LCMS. Upon completion, the reaction was quenched with sat. aq. NH₄Cl (add slowly to avoid strong gas evolution and exotherm). Sat. aq. NaHCO₃ was then added to avoid an emulsion, and the basic organic mixture was extracted with ether. The combined ether layers were washed with sat. aq. NaHCO₃, water, brine, dried Na₂SO₄, filtered, and concentrated to dryness to give the crude product. The crude product was purified by eluting through silica with EtOAc/Hexane (1:4 v/v). The purified product contained some residual 3-bromopropyl phthalimide, that did not interfere with subsequent synthetic steps. The material was taken on and used without further purification. Assume a quantitative yield.

Preparation of tert-butyl 1-(3-aminopropyl)-4-(2,4-dichlorophenyl)pyrrole-3-carboxylate (5).

The Pthalimido Pyrrole 4 (1 eq) was dissolved in ethanol and hydrazine (3 eq) at room temperature under nitrogen. Upon heating to reflux, the reaction generated a white precipitate. Stir at reflux until complete (~2 h) by TLC (CH₂Cl₂/acetonitrile (95:5 $v\dot{v}v$), the UV active product was at R_f= 0.2) and LCMS. Upon reaching completion, the reaction was allowed to cool to room temperature and the precipitate was vacuum-filtered off using a medium to fine cintered-glass filter. The filtrate was concentrated under reduced pressure to a gummy solid.

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The crude material was taken up in ethanol/EtOAc (1:1 v/v), stirred and the precipitate was filtered off in the same fashion as before. The filtrate was concentrated under reduced pressure and than dried in vacuo for 10-15 min. This process of adding ethanol/EtOAc, filtering and concentrating was done one more time or as needed to remove the majority of the white precipitate and residual hydrazine. The product was then dried in vacuo overnight. The material was used without further purification. Once dried, the reaction yielded the product as a glass (~87% yield over 2 steps).

Preparation of *tert*-butyl 1-{3-[(6-amino-5-nitro(2-pyridyl))amino]propyl}-4-(2,4-dichlorophenyl)pyrrole-3-carboxylate (7).

To the premixed dry reagents, pyrrole 5 (1 eq) and powdered 6-chloro-3-nitro-2-pyridylamine (6) (1.1 eq), was added the DMA followed by Hünig's base (2 eq) sequentially with stirring at rt. The reaction was then heated to 80 °C overnight. The reaction was followed by TLC (EtoAc/hexane (1:1 v/v), the UV active yellow product was at R_f = 0.25), HPLC and LCMS. Upon completion as judged by HPLC, the reaction was allowed to cool to 70 °C. Ethylene diamine (anhydrous) was then added to the reaction to destroy any remaining unreacted chloropyridine 6. After 15 min stirring at 70 °C, the reaction was cooled and quenched with the addition of sat. aq. NaHCO₃. The aqueous mixture was extracted with EtoAc, and the combined organic layers were washed with sat. aq. NaHCO₃, water, brine, dried, filtered, and concentrated to dryness to give the crude product as a brown-yellow solid. The crude product was purified by flash chromatography eluted with EtoAc/hexane (4:6 v/v). The purified SnAr adduct 7 was isolated in 58% yield as a yellow solid.

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Preparation of 1-{3-[(6-amino-5-nitro(2-pyridyl))amino]propyl}-4-(2,4-dichlorophenyl)pyrrole-3-carboxylic acid (8).

In a vial, TFA (catalytic amount) was added to a stirred mixture of tert-butyl ester pyrrole 7 (1 eq), water (.1%), and CH_2Cl_2 at rt. The vial stirred at room temperature until done (~12 h. The reaction was then concentrated under reduced pressure at room temperature and dried in vacuo. The crude residue was dissolved again in CH_2Cl_2 and concentrated under reduced pressure at rt. The material was used in the final coupling step without further purification as the TFA salt

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Preparation of N-((1S)-2-hydroxy-isopropyl)(1-{3-[(6-amino-5-nitro(2-pyridyl))amino]propyl}-4-(2,4-dichlorophenyl)pyrrol-3-yl)carboxamide (9,).

(2S)-(+)-2-Aminopropan-1-ol (1.5 eq) was added to a stirred mixture of acid (8) (1 eq), HBTU (1.5 eq), Hünig's base (2 eq) and DMF (premixed sequentially in this order in a vial) at room temperature under argon. The reaction was stirred for 3-4 h until complete as shown by LCMS and HPLC. The reaction mixture was subsequently diluted with EtOAc, washed with NaHCO₃ and concentrated to afford a powder in a 70% yield.

Nomenclature for the Example compounds was provided using ACD Name version 5.07 software (November 14, 2001) available from Advanced Chemistry Development, Inc. Some of the compounds and starting materials were named using standard IUPAC nomenclature.

The compounds of Table 34 were synthesized following the synthetic methodology described above in the Examples and Schemes, and screened following methods 1 and 2 below. The precursors are readily recognizable by one skilled in the art and are commercially available from Aldrich (Milwaukee, WI) or Acros Organics (Pittsburgh, PA), among others.

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Screening methods for SMIP/SMIS compounds

Method 1

Candidate small molecule immuno-potentiators can be identified *in vitro*. Compounds are screened *in vitro* for their ability to activate immune cells. One marker of such activation is the induction of cytokine production, for example TNF- α production. Apoptosis inducing small

molecules may be identified having this activity. These small molecule immuno-potentiators have potential utility as adjuvants and immuno-therapeutics.

In an assay procedure (High Throughput Screening (HTS)) for small molecule immune potentiators (SMIPs), human peripheral blood mononuclear cells (PBMC), 500,000 per mL in RPMI 1640 medium with 10% FCS, were distributed in 96 well plates (100,000 per well) already containing 5μ M of compound in DMSO. The PBMCs were incubated for 18 h at 37°C in 5% CO₂. Their ability to produce cytokines in response to the small molecule compounds is determined using a modified sandwich ELISA.

Briefly supernatants from the PBMC cultures were assayed for secreted TNF using a primary plate bound antibody for capture followed by a secondary biotinylated anti-TNF antibody forming a sandwich. The biotinylated second antibody was then detected using streptavidin-Europium and the amount of bound europium was determined by time resolved fluorescence. SMIP compounds were confirmed by their TNF inducing activity that was measured in the assay as increased Europim counts over cells incubated in RPMI medium alone. "Hits" were selected based on their TNF-inducing activity relative to an optimal dose of lipopolysaccaride LPS (1 μ g/ml), a strong TNF inducer. The robustness of the assay and low backgrounds allowed for the routine selection of hits with ~10% of LPS activity that was normally between 5-10X background (cells alone). Selected hits are then subjected to confirmation for their ability to induce cytokines from multiple donors at decreasing concentrations. Those compounds with consistent activity at or below 5μ M are considered confirmed for the purposes of this assay. The assay is readily modified for screening for compounds effective at higher or lower concentrations.

Method 2

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Each of the compounds in the above Table 34 elicited TNF- α production in human peripheral blood mononuclear cells. Many of the compounds showed activity at less than 20 μ M with respect to production of TNF- α . Many of these compounds showed activity at less than 5 μ M with respect to production of TNF- α . Many of these compounds showed activity in the production of TNF- α at less than 1.5 μ M.

For this reason, each of the R groups of any of the compounds listed in Table 34_are preferred. Additionally, because of the excellent activity of each of the compounds, each of these compounds is individually preferred and is preferred as a member of a group that includes any or all of the other compounds and each compound is preferred in methods of modulating

immunopotentiation and in methods of treating biological conditions associated therewith, for example to be used as a vaccine adjuvant. Each of the compounds is also preferred for use in preparation of medicaments for vaccines, immunopotentiation, reducing tumor growth and in treating biological conditions mediated therefrom.

In addition to the procedure described above, methods of measuring other cytokines (e.g. IL1-beta, IL-12, IL-6, IFN-gamma, IL-10 etc.) are well known in the art and can be used to find active SMIP compounds of the present invention.

Compounds may be useful that cause production of TNF-a at higher concentrations, such as 100 µM, 200 µM or 300 µM in the assays described herein. For example Loxoribine causes useful production of TNF-a at 300µM (see Pope et al Immunostimulatory Compound 7-Allyl-8-Oxoguanosine (Loxoribine) Induces a Distinct Subset of Murine Cytokines Cellular Immunology 162: 333-339 (1995)).

The subject invention also includes isotopically-labeled antiviral compounds, that are structurally identical to those disclosed above, but for the fact that one or more atoms are replaced by an atom having an atomic mass or mass number different from the atomic mass or mass number usually found in nature. Examples of isotopes that can be incorporated into antiviral compounds of the invention include isotopes of hydrogen, carbon, nitrogen, oxygen, phosphorous, sulfur, fluorine and chlorine, such as ²H, ³H, ¹³C, ¹⁴C, ¹⁵N, ¹⁸O, ¹⁷O, ³¹P, ³²P, ³⁵S, ¹⁸F and ³⁶Cl, respectively. Antiviral compounds of the present invention, derivatives thereof, and pharmaceutically acceptable salts of said compounds and of said derivatives that contain the aforementioned isotopes and/or other isotopes of other atoms are within the scope of this invention. Certain isotopically-labeled antiviral compounds of the present invention, for example those into which radioactive isotopes such as ³H and ¹⁴C are incorporated, are useful in drug and/or substrate tissue distribution assays. Tritiated, i.e., ³H, and carbon-14, i.e., ¹⁴C, isotopes are particularly preferred for their ease of preparation and detectability. Further, substitution with heavier isotopes such as deuterium, i.e., ²H, may afford certain therapeutic advantages resulting from greater metabolic stability, for example increased in vivo half-life or reduced dosage requirements and, hence, may be preferred in some circumstances. Isotopically labeled antiviral compounds of this invention and derivatives thereof can generally be prepared by carrying out known or referenced procedures and by substituting a readily available isotopically labeled reagent for a non-isotopically labeled reagent. In accordance with the present invention, methods are provided for the administration of an

compositions comprising a SMIP compound, an antigen, and optionally other adjuvants.

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As adjuvants, the SMIP compounds are combined with antigens and delivery systems to form a final immunogenic composition or vaccine product.

As immunotherapeutics, the SMIP compounds are used alone or in combination with other therapies for treatment of SARS.

Those of ordinary skill in the art will recognize that physiologically active antiviral compounds, SMIPs or SMISs that have accessible hydroxy groups are frequently administered in the form of pharmaceutically acceptable esters. The antiviral compounds of this invention can be effectively administered as an ester, formed on the hydroxy groups, just as one skilled in pharmaceutical chemistry would expect. It is possible, as has long been known in pharmaceutical chemistry, to adjust the rate or duration of action of the antiviral compound by appropriate choices of ester groups.

Other compounds that can be used in combination with the therapeutic agents described herein include, pentoxifylline (PTX), methylprednisolone, trimetrexate (Neutrexin), Zadaxin (thymosin alpha 1), optionally substituted 5-aminomethinimino-3-methyl-4-isoxazolecarboxylic acid phenylamides, cyclosporine A (CsA), 6-oxo-1,4,5-thiadiazin[2,3-b]quinazoline, 3-amino-2(1H)-thioxo-4(3H)-quinazolinone, gangciclovir, glycyrrhizin, tetracyclines, aminoglycosides, quinolones, bicyclam (1,4-Bis(1,4,8,11-tetraazacyclotetradec-1-ylmethyl)benzene octahydrochloride dihydrate), rapamycin, wortmannin, enalapril, roquinimex/linomide, inactivin, DNCB, AG7088, 9-aminocamptothecin (CFT-11), loxorobine, bropirimine, Ononase @ (ranpirnase), statins, such as: lovastatin—Mevacor®, pravastatin—Pravachol®, simvastatin—Zocor®, fluvastatin—Lescol®, atorvastatin—Lipitor® and rosuvastatin—Crestor®.

As used herein, the term "effective amount" means an amount of antiviral compound of the compositions, kits and methods of the present invention that is capable of treating the symptoms of the described conditions. The specific dose of a compound administered according to this invention will, of course, be determined by the particular circumstances surrounding the case including, for example, the compound administered, the route of administration, the state of being of the patient, and the severity of the condition being treated.

The dose of an antiviral compound of this invention to be administered to a subject is rather widely variable and subject to the judgment of the attending physician. It should be noted that it may be necessary to adjust the dose of a compound when it is administered in the form of a salt, such as a laureate, the salt forming moiety of which has an appreciable molecular weight.

The following dosage amounts and other dosage amounts set forth elsewhere in this description are for an average human subject having a weight of about 65 kg to about 70 kg. The skilled practitioner will readily be able to determine the dosage amount required for a subject whose weight falls outside the 65 kg to 70 kg range, based upon the medical history of the

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subject and the presence of diseases, e.g., diabetes, in the subject. Calculation of the dosage amount for other forms of the free base form such as salts or hydrates is easily accomplished by performing a simple ratio relative to the molecular weights of the species involved.

In general, the pharmaceutical compositions will include at least one antiviral compound in combination with a pharmaceutically acceptable vehicle, such as saline, buffered saline, 5% dextrose in water, borate-buffered saline containing trace metals or the like. Formulations may further include one or more excipients, preservatives, solubilizers, buffering agents, lubricants, fillers, stabilizers, etc. Methods of formulation are well known in the art and are disclosed, for example, in "Remington's Pharmaceutical Sciences," Mack Pub. Co., New Jersey (1991) or "Remington: The Science and Practice of Pharmacy," 20th ed., Lippincott Williams & Wilkins, Baltimore, Maryland (2000), incorporated herein by reference.

Pharmaceutical compositions for use within the present invention can be in the form of sterile, non-pyrogenic liquid solutions or suspensions, coated capsules, suppositories, lyophilized powders, transdermal patches or other forms known in the art.

Many of the active ingredient antiviral compounds are known to be absorbed from the alimentary tract, and so it is usually preferred to administer a compound orally for reasons of convenience. However, the compounds may equally effectively be administered intravenously, subcutaneously, percutaneously, or as suppositories for absorption by the rectum or vagina, if desired in a given instance. All of the usual types of compositions may be used, including tablets, chewable tablets, capsules, solutions, parenteral solutions, troches, suppositories and suspensions. Compositions are formulated to contain a daily dose, or a convenient fraction of daily dose, in a dosage unit, that may be a single tablet or capsule or convenient volume of a liquid.

Capsules are prepared by mixing the compound or compounds with a suitable diluent and filling the proper amount of the mixture in capsules. The usual diluents include inert powdered substances such as starch of many different kinds, powdered cellulose, especially crystalline and microcrystalline cellulose, sugars such as fructose, mannitol and sucrose, grain flours and similar edible powders.

Tablets are prepared by direct compression, by wet granulation, or by dry granulation.

Their formulations usually incorporate diluents, binders, lubricants and disintegrators as well as the compound or compounds. Typical diluents include, for example, various types of starch, lactose, mannitol, kaolin, calcium phosphate or sulfate, inorganic salts such as sodium chloride and powdered sugar. Powdered cellulose derivatives are also useful. Typical tablet binders are substances such as starch, gelatin and sugars such as lactose, fructose, glucose and the like.

Natural and synthetic gums are also convenient, including acacia, alginates, methylcellulose,

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polyvinylpyrrolidine and the like. Polyethylene glycol, ethylcellulose and waxes can also serve as binders.

A lubricant is generally necessary in a tablet formulation to prevent the tablet and punches from sticking in the die. The lubricant is chosen from such slippery solids as talc, magnesium and calcium stearate, stearic acid and hydrogenated vegetable oils.

Tablet disintegrators are substances that swell when wetted to break up the tablet and release the compound or compounds. They include starches, clays, celluloses, algins and gums, more particularly, corn and potato starches, methylcellulose, agar, bentonite, wood cellulose, powdered natural sponge, cation-exchange resins, alginic acid, guar gum, citrus pulp and carboxymethylcellulose, for example, may be used as well as sodium lauryl sulfate.

Tablets are often coated with sugar as a flavor and sealant, or with film-forming protecting agents to modify the dissolution properties of the tablet. The compounds may also be formulated as chewable tablets, by using relatively large amounts of pleasant-tasting substances such as mannitol in the formulation, as is now well-established in the art.

When it is desired to administer a compound as a suppository, the typical bases may be used. Cocoa butter is a traditional suppository base, that may be modified by addition of waxes to raise its melting point slightly. Water-miscible suppository bases comprising, particularly, polyethylene glycols of various molecular weights are in wide use.

The effect of the compounds may be delayed or prolonged by proper formulation. For example, a slowly soluble pellet of the compound may be prepared and incorporated in a tablet or capsule. The technique may be improved by making pellets of several different dissolution rates and filling capsules with a mixture of the pellets. Tablets or capsules may be coated with a film that resists dissolution for a predictable period of time. Even the parenteral preparations may be made long-acting by dissolving or suspending the compound or compounds in oily or emulsified vehicles that allow dispersion slowly in the serum.

The combinations of this invention may be administered in a controlled release formulation such as a slow release or a fast release formulation. Such controlled release formulations of the combination of this invention may be prepared using methods well known to those skilled in the art. The method of administration will be determined by the attendant physician or other person skilled in the art after an evaluation of the subject's condition and requirements.

The term "prodrug" means compounds that are transformed *in vivo* to yield an antiviral compound of the present invention. The transformation may occur by various mechanisms, such as through hydrolysis in blood. A good discussion of the use of prodrugs is provided by T. Higuchi and W. Stella, "Pro-drugs as Novel Delivery Systems," Vol. 14 of the A.C.S. Symposium Series, and in Bioreversible Carriers in Drug Design, ed. Edward B. Roche, American Pharmaceutical Association and Pergamon Press, 1987. The term, "prodrug" also

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encompasses mutual prodrugs in which one or more antiviral compounds are combined in a single molecule that may then undergo transformation to yield the individual antiviral compounds of the present invention.

For example, if an antiviral compound of the present invention contains a carboxylic acid functional group, a prodrug can comprise an ester formed by the replacement of the hydrogen atom of the acid group with a group such as (C_1-C_8) alkyl, (C_2-C_{12}) alkanoyloxymethyl, 1-(alkanoyloxy)ethyl having from 4 to 9 carbon atoms, 1-methyl-1-(alkanoyloxy)-ethyl having from 5 to 10 carbon atoms, alkoxycarbonyloxymethyl having from 3 to 6 carbon atoms, 1-(alkoxycarbonyloxy)ethyl having from 4 to 7 carbon atoms, 1-methyl-1-

(alkoxycarbonyloxy)ethyl having from 5 to 8 carbon atoms, N-(alkoxycarbonyl)aminomethyl having from 3 to 9 carbon atoms, 1-(N-(alkoxycarbonyl)amino)ethyl having from 4 to 10 carbon atoms, 3-phthalidyl, 4-crotonolactonyl, gamma-butyrolacton-4-yl, di-N,N-(C₁-C₂)alkylamino(C₂-C₃)alkyl (such as β -dimethylaminoethyl), carbamoyl-(C₁-C₂)alkyl, N,N-di(C₁-C₂)alkylcarbamoyl-(C₁-C₂)alkyl and piperidino-, pyrrolidino- or morpholino(C₂-C₃)alkyl.

Similarly, if an antiviral compound of the present invention comprises an alcohol functional group, a prodrug can be formed by the replacement of the hydrogen atom of the alcohol group with a group such as $(C_1\text{-}C_6)$ alkanoyloxymethyl, 1-($(C_1\text{-}C_6)$ alkanoyloxy)ethyl, 1-methyl-1-($(C_1\text{-}C_6)$ alkanoyloxy)ethyl, $(C_1\text{-}C_6)$ alkoxycarbonyloxymethyl, N-($C_1\text{-}C_6$)alkoxycarbonylaminomethyl, succinoyl, $(C_1\text{-}C_6)$ alkanoyl, α -amino $(C_1\text{-}C_4)$ alkanoyl, arylacyl and α -aminoacyl, or α -aminoacyl- α -aminoacyl, where each α -aminoacyl group is independently selected from the naturally occurring L-amino acids, $P(O)(OH)_2$, $P(O)(O(C_1\text{-}C_6)$ alkyl)2 or glycosyl (the radical resulting from the removal of a hydroxyl group of the hemiacetal form of a carbohydrate).

If an antiviral compound of the present invention comprises an amine functional group, a prodrug can be formed by the replacement of a hydrogen atom in the amine group with a group such as R^X -carbonyl, R^X O-carbonyl, NR^XR^X -carbonyl where R^X and R^X - are each independently ((C₁-C₁₀)alkyl, (C₃-C₇)cycloalkyl, benzyl, or R^X -carbonyl is a natural α -aminoacyl or natural α -aminoacyl-natural α -aminoacyl, -C(OH)C(O)OYX wherein (YX is H, (C₁-C₆)alkyl or benzyl), -C(OYXO) YXI wherein YXO is (C₁-C₄) alkyl and YXI is ((C₁-C₆)alkyl, carboxy(C₁-C₆)alkyl, amino(C₁-C₄)alkyl or mono-N- or di-N,N-(C₁-C₆)alkylaminoalkyl, -C(YXO) YXI wherein YXO is H or methyl and YXI is mono-N- or di-N,N-(C₁-C₆)alkylamino, morpholino, piperidin-1-yl or pytrolidin-1-yl.

The compositions for use in accordance with the present invention may be formulated in a conventional manner using one or more physiologically acceptable carriers or excipients.

Antiviral, SMIP, SMIS, or other immunomodulating compounds are prepared or obtained as described herein and in the US Patents and published international patent applications listed in

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Table 1, Table 2, Table 34 and Table 35. The antiviral compounds can be formulated in pharmaceutically acceptable compositions suitable for delivery to the lungs. Particular formulations include dry powders, liquid solutions or suspensions suitable for nebulization and propellant formulations suitable for use in metered dose inhalers. The preparation of such formulations is well know to those skilled in the art, and is described in US Patent Nos. 5,814,607 and 5,654,007 and in the US Patents and published international patent applications listed in Table 3 the disclosures of which are incorporated herein by reference.

Dry powder formulations will comprise an antiviral compound in a dry, optionally lyophilized form with a particle size within a preferred range for deposition within the lung. Typically the particle size for deposition in the lung will range between 1 and 5 μ m. When systemic delivery of the antiviral compound via absorption from the lung into the bloodstream is desired the antiviral compound formulation particle size is generally between 0.1 and 2 μm in size. The preferred size range of particles can be produced using methods such as jet-milling, spray drying and solvent precipitation, for example. Dry powder devices typically require a powder mass in the range from about 1 mg to 100 mg to produce an aerosolized dose. Thus, the antiviral compound will typically be combined with a pharmaceutically acceptable dry bulking powder. Preferred dry bulking powders include sucrose, lactose, trehalose, human serum albumin (HSA), phospholipids and glycine as well as those disclosed in the documents listed in Table 3. Dry powders can be administered to the subject in conventional dry powder inhalers. For liquid formulations the antiviral compound can be dissolved in any recognized physiologically acceptable carrier for use in delivery of aerosolized formulations. Such carriers include buffered and unbuffered aqueous solutions for water soluble compounds, and physiological solutions including saline solution (preferably between 0.2 and 2 N NaCl). For antiviral compounds with limited solubility, other liquid vehicles such as ethanol, propylene glycol and ethanol-propylene combinations may be used. The antiviral compounds may also be administered as solids in suspension.

For administration by inhalation, the compositions for use according to the present invention are conveniently delivered in the form of an aerosol spray administered via pressurized packs or a nebulizer, with the use of a propellant, e.g., air, dichlorordifluoromethane, dichloroterafluoroethane or other suitable gas. Preferably, for incorporation into the aerosol propellant, the antiviral compound formulations of the present invention will be processed into respirable particles as described above for the dry powder formulations. The particles are then suspended in the propellant, optionally being coated with a surfactant to enhance their disbursement. In the use of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount.

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Commercially available jet nebulizers are available and may be used to deliver aerosolized antiviral compound to a subject. Such jet nebulizers include, but are not limited to, those supplied by AeroTech 11 (CIS-US, Bedford, Mass.). In addition, for delivery of acrosolized antiviral compound to the lungs of a subject an oxygen source can be attached to the nebulizer providing a flow rate of, for example, 10 L/min. In general, inhalation is performed over a 5-40 minute time interval through a mouthpiece during spontaneous respiration. The present invention provides for novel compositions comprising a suitable carrier and aerosolized antiviral compound in doses sufficient to reduce or ameliorate viral load and SARS symptoms in subjects having SARS. Such doses can be lower than corresponding systemic doses that may be used to those generally used to reduce or ameliorate viral load and SARS symptoms in subjects having SARS.

The antiviral, SMIP, SMIS, and immunomodulating compositions of the present invention may be administered with a steroidal anti-inflammatory drug for the treatment of SARS and SARS symptoms. Examples of steroidal anti-inflammatory drugs of the invention include hydrocortisone, prednisolone, dexamethasone, triamcinolone acetonide, fluorinolone acetonide, fluorinolone acetate, betamethasone, etc.

The antiviral compound composition of the invention is nebulized predominantly into particle sizes allowing a delivery of the drug into the terminal and respiratory bronchioles. For efficacious delivery of antiviral compound to the lung endobronchial space of airways in an aerosol, the formation of aerosol particles having mass medium average diameter predominantly between 1 to 5 μ m is necessary. The formulation must additionally provide conditions that would not adversely affect the functionality of the airways. Consequently, the formulation must contain enough of the drug formulated under the conditions that allow its efficacious delivery while avoiding undesirable reaction.

For liquid solutions and suspensions, the choice of the nebulizer is made from among commercially available nebulizers. The jet nebulizers known as Sidestream O, obtained from Medicaid and Pari LCS, LC Plus, and eFlow obtained from Pari Respiratory Equipment, Richmond, Virginia, are examples of typical nebulizers suitable for the practice of the invention. Ultrasonic nebulizers that produce appropriate particle sizes of about 1 to 5 μ m such as Aerosonic by DeVilbiss and UltraAire by Omron are also suitable.

Advantageously, the present invention also provides for a kit for use by a consumer for the treatment and/or prevention of SARS. Such a kit comprises: (a) a pharmaceutical composition comprising a therapeutically effective amount of at least one compound from among those described herein or listed in Table 34 and Table 35 or described in the US Patents and published international patent applications listed in Table 1, Table 2, and Table 35 and a pharmaceutically acceptable carrier, vehicle or diluent; (b) a container for holding the pharmaceutical composition;

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and, optionally, (c) instructions describing a method of using the pharmaceutical compositions for the treatment and or the prevention of SARS. The kit may optionally contain a plurality of antiviral compounds for the treatment of SARS wherein the anti-viral compounds are selected from 3C-like protease inhibitors and papain-like protease inhibitors. In a further embodiment, the kit contains an antiviral compound which is an RNA-dependent RNA polymerase inhibitor. When the kit comprises more than one antiviral compound, the antiviral compounds contained in the kit may be optionally combined in the same pharmaceutical composition.

A "kit" as used in the instant application includes a container for containing the separate compositions such as a divided bottle or a divided foil packet. The container can be in any conventional shape or form as known in the art that is made of a pharmaceutically acceptable material, for example a paper or cardboard box, a glass or plastic bottle or jar, a re-sealable bag (for example, to hold a "refill" of tablets for placement into a different container), or a blister pack with individual doses for pressing out of the pack according to a therapeutic schedule. The container employed can depend on the exact dosage form involved, for example a conventional cardboard box would not generally be used to hold a liquid suspension. It is feasible that more than one container can be used together in a single package to market a single dosage form. For example, tablets may be contained in a bottle that is in turn contained within a box.

An example of such a kit is a so-called blister pack. Blister packs are well known in the packaging industry and are being widely used for the packaging of pharmaceutical unit dosage forms (tablets, capsules, and the like). Blister packs generally consist of a sheet of relatively stiff material covered with a foil of a preferably transparent plastic material. During the packaging process, recesses are formed in the plastic foil. The recesses have the size and shape of individual tablets or capsules to be packed or may have the size and shape to accommodate multiple tablets and/or capsules to be packed. Next, the tablets or capsules are placed in the recesses accordingly and the sheet of relatively stiff material is sealed against the plastic foil at the face of the foil that is opposite from the direction in which the recesses were formed. As a result, the tablets or capsules are individually sealed or collectively sealed, as desired, in the recesses between the plastic foil and the sheet. Preferably the strength of the sheet is such that the tablets or capsules can be removed from the blister pack by manually applying pressure on the recesses whereby an opening is formed in the sheet at the place of the recess. The tablet or capsule can then be removed via said opening.

It maybe desirable to provide a written memory aid, where the written memory aid is of the type containing information and/or instructions for the physician, pharmacist or subject, e.g., in the form of numbers next to the tablets or capsules whereby the numbers correspond with the days of the regimen which the tablets or capsules so specified should be ingested or a card that contains the same type of information. Another example of such a memory aid is a calendar

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printed on the card e.g., as follows "First Week, Monday, Tuesday," . . . etc "Second Week, Monday, Tuesday, . . . " etc. Other variations of memory aids will be readily apparent. A "daily dose" can be a single tablet or capsule or several tablets or capsules to be taken on a given day. Also a daily dose of one or more component(s) of the kit can consist of one tablet or capsule while a daily dose of another one or more component(s) of the kit can consist of several tablets or capsules.

Another specific embodiment of a kit is a dispenser designed to dispense the daily doses one at a time in the order of their intended use. Preferably, the dispenser is equipped with a memory-aid, so as to further facilitate compliance with the regimen. An example of such a memory-aid is a mechanical counter that indicates the number of daily doses that has been dispensed. Another example of such a memory-aid is a battery-powered micro-chip memory coupled with a liquid crystal readout, or audible reminder signal which, for example, reads out the date that the last daily dose has been taken and/or reminds one when the next dose is to be taken.

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EXAMPLES

Example 1-EXAMPLE of a SARS VIRUS ISOLATE

A SARS virus was isolated from clinical specimens of a patient in Frankfurt, Germany (FRA). The isolate was grown in Vero cells. RNA of the SARS virus was extracted and amplified by RT-PCR. Nucleotide sequence of the viral genome was determined by direct sequencing of the PCR product. Computer analysis was used to predict the features of the genome, to compare it to previously known coronaviruses and to the sequence of different SARS virus isolates.

More specifically, isolation and sequence was performed as follows. After the third passage of the SARS virus in Vero cells, viral particles were purified by ultra centrifugation from 3×10^7 cells supernatant. Viral RNA was extracted by Triazol method (Gibco-BRL). Viral RNA (200 ng) was transcribed into cDNA with avian RNaseH- thermostable reverse transcriptase following the instructions of the manufacturer (ThermoScript RT System, Invitrogen). Briefly, either 50 pmoles of oligo (dT)₂₀ (SEQ ID NO: 7389) or 25 ng of random hexamers were used to prime the RT reaction in a 20 μ l final volume. Amplification and sequencing of the SARS genome were accomplished by direct sequencing of PCR products obtained with: i) specific primers from conserved regions of homology found through multiple alignment among known coronaviruses; ii) oligonucleotides designed around short sequences of SARS isolates available on the Web through WHO network laboratories; iii) degenerate primers to amplify the cDNA mixture with multiple overlapping fragments as end products. Gap closure

was realized by long distance PCR with high fidelity Taq (Expand High Fidelity system, Roche) using primers designed on selected fragments. Sequence was collected by primer walking using a BigDye terminator chemistry (Applied Biosystems) and an automated DNA sequencer (3700 capillary model, Applied Biosystems). After obtaining a first pass of the entire genome, a set of both forward and reverse primers were used to amplify and sequence *de novo* the genome using as a template DNA segments of 2 kb on average. Readings from overlapping fragments were automatically assembled by AutoAssembler (Applied Biosystems) and the 29,740 bp contiguous edited manually.

Computer analysis of the sequence was performed as follows. The GCG Wisconsin Package suite (version 10.0) was used for computer analysis of gene and protein sequences. The PSORT program (http://psort.nibb.ac.jp/) was used for localization predictions. For secondary structure analysis, the PHD software available on the Web at http://cubic.bioc.columbia.edu/predictprotein/ was applied. The PSI-BLAST algorithm was used for homology searches (http://www.ncbi.nlm.nih.gov/blast) using the non-redundant protein database. ClustalW was applied to obtain multiple sequence alignments of gene and protein sequences. The LearnCoil-VMF program was used to predict coiled-coil regions in the spike proteins (http://learncoil-wnf.lcs.mit.edu/cgi-bin/vnf). Leucine zippers were predicted with the program 2ZIP, available at http://2Zip.molgen.mpg.de.

Phylogenetic analysis was performed using the neighbor-joining algorithm as implemented in the program NEIGHBOR within the Phylogeny Inference Package (Phylip) (Felsenstein J 1993, program distributed by the author). Bootstrap analysis was always performed with 100 replicates using the program Seqboot. Trees were handled and displayed using TreeView. The program HMMER was used to generate sequence profiles from multiple sequence alignments of the S1 domains of spike proteins. Subsequently, the HMMPFAM program was used to compare the S1 domain of SARS spike to the profiles.

The genome of this SARS virus isolate is 29,740 bases long and the overall structure of the genome is similar to that of the three known groups of coronaviruses. Starting from the 5' end a leader sequence, an untranslated region (UTR) and two overlapping open reading frames coding for one polyprotein containing the enzymes necessary for replication can be identified. They are followed by a region coding for the spike (S), envelope (E), matrix (M), nucleocapsid (N) structural proteins and eight additional ORFs specific for the SARS virus. At the 3'-end of the genome a UTR with a poly(A) is located. The overall homology to coronaviruses groups 1, 2 and 3 is low and therefore the SARS virus belongs to a new group (group 4) of coronavirus. More detailed analysis of the spike protein amino acid sequence shows that the SARS virus isolate is more closely related to coronavirus group 2.

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The complete genome sequence of the SARS virus isolate is 29,740 bp in length. The sequence is available on Genbank and has a GC content of 40.8%, comparable with that of known viruses of the same family. Genome structure is similar to that of other coronaviruses. 14 open reading frames have been predicted. The principal features of the genome and gene products are illustrated reported in Figure 17 and Table 10. The comparison between the SARS genome and those of group 1, 2 and 3 coronaviruses is reported in Figure 18.

Nucleotides 1-73 contain a predicted RNA leader sequence followed by an untranslated region (UTR) of 197 nucleotides. The UTR is followed by two overlapping open reading frames (ORF1a, ORF1b), which encompass two-thirds of the genome (nucleotides 265-21485). They encode for a large polyprotein, which is predicted to be processed by viral proteases to generate the replicase complex. The 3' part of the genome contains the genes coding for the four structural proteins (S, spike protein, E, envelope protein, M, matrix glycoprotein, and N, nucleocapsid protein), and eight predicted ORFs of unknown function (Figure 17). Finally, at the 3' end of the genome, we found a second UTR of 340 bases followed by a poly(A) tract. We identified a putative intergenic (IG) sequence also referred to as transcription-associated sequence (TAS), which is a typical feature for coronaviruses. The IG sequence is characterized by 6-18 nucleotides present at the 3' end of the leader and can be found in front of each gene. The IG sequence plays a key role in RNA transcription and its regulation. The IG sequence of the SARS virus is characterized by the sequence SEQ ID NO: 7293 and is present nine times in the genome (Figure 17). The sequence of the leader and IG are peculiar for each coronavirus and represent a specific signature for the virus.

The Replicase Region

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The replicase gene, ORF1ab (SEQ ID NO: 7232), consists of two overlapping ORFs, ORF1a and ORF1b, which can be translated as a single polyprotein by frame shift of the ribosome in position 13,393, within the polymerase encoding region. See Brierley et al, *Embo J* 1987: 6(12): 3779-3785. As expected, a stem-loop sequence is present ten base pairs downstream of this site (SEQ ID NO: 7390;

5'-CGGTGTAAGTGCAGCCCGTCTTACACCG-3'). The polyprotein is cleaved co- and/or post-translationally into multiple proteins by its own encoded proteases. Using the cleavage consensus sequence and by analogy with other coronaviruses, we have mapped the possible cleavage sites of the polyprotein and have identified 14 products, which comprise the leader protein p28, the homologue of the MHV p65 protein and other twelve proteins, named from nsp1 to nsp13 (nsp, non structural protein) (Figure 17 and Table 10). The amino acid sequence analysis suggests the presence of several functional motifs within the putative ORF1ab proteins. In particular, we have mapped two potential proteases (nsp1 and nsp2), one growth factor-like motif (nsp7) within ORF1a, whereas in ORF1b we identified the RNA polymerase (nsp9), and a

predicted helicase (nsp10). The other predicted cleavage products (nsp3, nsp4, nsp5, nsp6, nsp11, nsp12 and nsp13) are proteins of unknown function. Many of these proteins are presumably present in the RNA replication complex, which is associated with the membranous structures in the infected cells. In particular, nsp3 and nsp4 contain hydrophobic domains. As shown in Figure 18, the replicase region of SARS has a similar organization to group 1, 2 and 3 coronaviruses; however, the overall aminoacid conservation is low (Table 11). The most conserved proteins are the polymerase and the helices.

Nsp1 is the papain-like cysteine protease (PLP), which cleaves the first two protein products (leader protein p28 and p65 homologue). Within the nsp1 of MHV, two domains with papain-like protease activity (PLP1 and PLP2) have been mapped, (Kanjanahaluethai et al (2000) J. Virol 74(17):7911-21) which are also conserved with Bovine, transmittable gastroenteritis virus (TGV) and Human 229E coronaviruses. However, by sequence alignment with the SARS nsp1, we identified only one PLP domain containing the catalytic residues Cys833 and His994.

Nsp2 is the chymotrypsin-picomavirus 3C-like protease (3CLp), which is responsible for the post-translational processing of the other 12 proteins, most of them cleaved at Q/A or Q/S sites. (Ziebuhr et al (1999) J. Virol 73(1):177-85). It also performs autoproteolytic activity. The principal catalytic residues are well conserved with other coronaviruses and are located at position His41 and Cys145. Furthermore, even the conserved aminoacids Tyr161 and His163, which are believed to be involved in substrate recognition and to be indispensable for proteolytic activity, (Hegyi et al (2002) J. Gen Virol 83(Pt3):581-593) were found in the sequence of the SARS 3CLD.

The invention includes the orf1ab sequence of SEQ ID NO: 9960 and the orf1a sequence of SEQ ID NO: 9961, including fragments, variants, homologs, etc. thereof.

25 The Structural Region

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Analysis of the nucleotide sequence at the 3' part of the SARS genome identified 12 predicted open reading frames. They are coded within 8.2 kb and comprise the four structural proteins S, E, M and N, common for all coronaviruses and eight predicted ORFs, which are specific for this virus (Figure 18). SARS-specific IG sequences upstream of most ORFs (Figures 17 & 18) suggest that most genes are likely to be transcribed independently. Interestingly, sequences identical to the group 2 IG are also present at the end of the RNA leader and in front of the Matrix encoding gene and of ORF 10.

The spike is a type I glycoprotein, which forms the large spikes on the surface of the virion and is responsible for receptor-binding and membrane fusion. (Gallagher (2001) Adv Exp Med Biol 494: 183-92). The protein is 1255 residues long with 17 predicted N-glycosylation sites. It has a 13aa leader peptide and a 17 aa C-terminal membrane anchoring sequence (1202-1218).

Some (MHV, HCoV-OC43, AIBV and BCoV), but not all (TGV, FIPV, HCoV-229E) coronavirus spike proteins are proteolytically cleaved in two subunits, S1 and S2. S1 is supposed to form the bulbous head, which stays non-covalently linked to the C-terminal membrane anchor. Cleavage is mediated by a basic aminoacid sequence, which resembles the consensus sequence for a furin cleavage site. (Garten et al., Biochimie 1994; 76(3-4): 217-225). However, in case of this SARS virus isolate, we were not able to identify such a sequence, implicating that the S protein of this SARS virus isolate is unlikely to be cleaved during maturation. Secondary structure predictions indicated that the global architecture of the spike protein is conserved within all known coronaviruses. The S1 domain is mainly formed by beta sheets and likely adopts a globular fold, while in the S2 domain extensive alpha helical regions are predicted. In addition, the LearnCoil-VMF program, specifically designed to identify coiledcoil-like regions in viral membrane-fusion proteins, predicts two coiled-coils within S2, spanning aminoacids 900-1005 and 1151-1185, respectively (Figure 19). Both coiled-coil regions contain a leucine-zipper motif, which is also present in the spikes of all coronaviruses. Leucine zippers are known to promote protein oligomerization; since the spike proteins of TGV and MHV form hetero-trimers, (Delmas et al., J Virol 1990; 64(11):5367-5375) (Godeke, et al., J Virology 2000; 74(3):1566-1571) it is conceivable that in SARS leucine zippers play a role in promoting and/or stabilizing a similar quaternary structure. The spike protein plays a major role in the biology of coronaviruses because the S1 domain contains the receptor-binding domain and the virus neutralizing epitopes, while the S2 domain is involved in the process of membrane fusion, which is essential for virus infectivity. As expected, multiple sequence alignment of different spike proteins showed a major degree of variability within the S1 domain, whereas S2 is more conserved.

The envelope protein E is a very short polypeptide of 76 aa, involved in the morphogenesis of the virion envelope. (Godet et al., Virology 1992; 188(2):666-675). Computer analysis predicts a long transmembrane domain close to the N-terminus and two N-glycosylation sites. The level of aminoacid similarity with other coronaviruses is very low and the best homology is with the small envelope protein of the transmissible gastroenteritis virus (TGV).

The matrix glycoprotein (M) is a 221-residue polypeptide with a predicted molecular weight of 25 kDa. Computer analysis predicts a topology consisting of a short aminoterminal ectodomain, three transmembrane segments and a carboxyl terminus located at the interior side of the viral envelope. In analogy with the matrix glycoprotein of TGV, that of the avian infective bronchitis virus (AIBV) and that of the hypervirulent MHV-2 strain the SARS M glycoprotein is N-glycosylated at the N-terminus. SARS M protein shows highest similarity to group 2 viruses (Table 11).

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Finally, the nucleocapsid protein N is a 397-residue-long phosphoprotein that interacts with viral genomic RNA to form the nucleocapsid. The level of conservation with other coronaviruses is low, ranging from 26,9% identity with the HCoV-229E to 37,4% identity to the Bovine coronavirus (BcoV) (Table 11). Epitope analysis of the nucleocapsid protein has been carried out (Li et al. (2003) Geno Prot & Bioinfo 1:198-206) in which the epitope site at the C terminus of the protein was located as SEQ ID NO: 7394 (amino acids 371-407 of SEQ ID NO: 6052).

In addition to the above fundamental proteins, many viruses express a set of other peptides, which are generally dispensable for viability, but can influence the infectivity potential of the virus. (de Haan et al., Virology 2002; 296(1):177-189). These proteins are generally conserved within members of the same serogroup, but differ profoundly among the groups. For this reason, they are generally referred to as group-specific proteins (Figure 11). Members of the group 1, represented here by HcoV-229E, have two group-specific genes located between the S and E genes and sometimes one or two ORFs downstream of the N gene, preceding the 3' UTR region of the genome. Viruses of the group 2, with MHV as prototype, have two group-specific genes (2a and HE) between ORF1b and S, as well as other two between S and E genes. Finally, the group 3 viruses, represented by the prototype AIBV, have two group-specific genes between S and E and other two between the M and N genes.

With the exception of the hemagglutinin esterase HE, for which hemagglutinating and acetyl-esterase enzymatic activities have been demonstrated, all the other group-specific ORFs encode proteins whose role has not yet been established.

Interestingly, the arrangement of specific genes in the SARS genome is peculiar and the predicted ORFs do not display any significant homology with ORFs present in the other coronaviruses, nor with any other known protein from different organisms. Like viruses of the group 1 and 3, SARS lacks the HE hemagglutinin and does not contain ORFs between the ORF1b and the S gene. Furthermore, two predicted ORFs (ORF3 and ORF4) are encoded in the region between S and E, and superimpose for most of their length. ORF3 has an IG sequence 2 bp upstream of the ATG start codon. In contrast to the other groups, SARS contains five predicted ORFs in the region between M and N genes. ORF7 is located 10 bases downstream of the stop codon of M gene, and has an IG sequence 155 nucleotides upstream from the ATG start codon. Similarly, ORF8 and ORF10 present an IG right upstream of their ATG start codons. On the other hand, the 5' ends of ORF9 and ORF11 shortly superimpose with the flanking genes, and for this reason they do not need an IG to activate transcription. ORF12 totally superimposes with the N gene and shares very low homology with a 22kDa protein of the MHV virus, coded in the corresponding region.

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Despite the absence of indications of possible localization and function deriving from sequence similarity, ORF3, ORF7 and ORF8 contain hydrophobic segments, suggesting association with membrane structures. In addition, ORF3, the longest among the SARS specific gene, is the only one that encodes for a peptide containing a high number of predicted O-glycosylation sites (Table 11). Predicted N-glycosylation sites have been identified in ORF3, ORF11 and ORF12.

Two shorter ORFs in the non-structural regions are SEQ ID NOS: 9965 and 9966. The invention includes polypeptides with these sequences, and also fragments, variants, etc.

Phylogenetic analysis

The substitution frequency within 922 conserved bases from the pol gene of eleven coronaviruses from the three different serogroups has been used in the past to show that the variability within members of each serogroup is much smaller than between members of different serogroups, confirming the previously described serological groupings. (Stephensen et al., Virus Res 1999; 60(2):181-9). We used the 922 bp region of the pol gene of SARS and aligned it with the same fragment from other 12 coronaviruses. The tree obtained showed that the SARS virus is distinct from the other three groups of coronaviruses (Figure 20). Similar results were obtained using the full-length aminoacid sequences of pol, 3CL-protease and helicase from the replicase region and those of the spike and the matrix glycoproteins from the structural region (data not shown). These data confirmed that the entire genome of the SARS virus clusters in a new group (group 4) of coronavirus.

To gain more resolution for possible evolutionary relationships we performed the analysis using consensus sequences of predicted domains of the proteins. In particular, we generated consensus sequences of the S1 domain of the spike protein from the group 1 and group 2 and then we compared them to the S1 domain of the SARS spike. No consensus could be generated from the group 3 since only the spike protein of AIBV is known. Interestingly, the tree constructed from the alignment of SARS S1 with the consensus generated from the two groups of spike proteins was different from that in Figure 20, and showed a much closer relationship between SARS and group 2 coronaviruses (Figure 21A). Further analysis showed that 19 out of the 20 cysteines present in the SARS S1 domain are spatially conserved with the group 2 consensus sequence, while only five are maintained either within the group 1 and group 3 sequences (Figure 21B). Given the fundamental role played by cysteines in protein folding, it is likely that the S1 domain of SARS and group 2 coronaviruses share a similar spatial organization.

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Sequence variability between SARS coronaviruses

We compared the FRA sequence to the four complete SARS genomes available on the Web. A total of 30 mutations were detected. Nine of these mutations were silent while 21 resulted in aminoacid substitutions (Table 12). Within ORF1a, three silent and seven productive mutations were detected. In ORF1b, there were five silent and three productive mutations. One of the productive mutations was caused by two nucleotide substitutions resulting in a single aminoacid change. Five changes were located in the spike protein, four of these were productive and one silent. Two productive mutations were in ORF3 and in the matrix glycoprotein M. One productive mutation each was in ORF10 and in the nucleocapsid protein N.

The overall difference between FRA and TOR2 was of nine nucleotides resulting in two silent mutations and seven aminoacid changes. The difference between FRA and Urbani is 12 nucleotides, which result in five silent mutations and seven aminoacid changes. For CUHK 16 nucleotides were different, five of which were silent mutations. For FRA and HKU 14 nucleotide changes resulted in four silent and nine productive mutations.

EXAMPLE 2 - Production, Inactivation and Purification of Whole SARS Virus Using MCS Chromatography Resin Purification Followed by Density Gradient Ultracentrifugation

A SARS isolate FRA1 (EMBL: AY310120) was passaged on VERO cells that were cultivated in DMEM (Gibco: Cat No. 21969-035, Lot No. 3078864), Penicillin/Strep (Gibco: Cat No. 15070-063, Lot No. 1120042), and 3% FCS (Gibco: Cat No. 10270-106, Lot No. 40F6130K) at 37°C, 5% CO₂. Trypsin (Gibco: Cat No. 25300-054, Lot No. 3078729) was used

for detaching the cells.

For virus production the third passage was used for inoculation of VERO cells at a moi of ~0.1. Cells were incubated with the virus for 1 h at 37°C in infection medium (DMEM without PS, FCS); after 1h cells were washed twice and further incubated at 37°C for 48 h in the presents.

of 3% FCS and antibiotics. The supernatant was harvested 48 hours post infection (p.i.) and precleared by centrifugation at 3000 rpm at 4°C for 10 min.

The SARS virus was inactivated by β -propiolactone (BPL) treatment (1:2000) for 18 h at 4°C, followed by 3 h at 37°C. Testing the virus on successful inactivation, VERO cells were incubated with 10 ml BPL treated supernatant for 4 days at 37°C; subsequentially, the supernatant was transferred to a fresh VERO cell culture and further incubated for another 4 days. Cells were checked for cytopathic effect (CPE).

200 ml of the BPL-inactivated SARS virus harvest was then clarified using a 0.65 µmpore-size filter (47 mm diameter) to pass virus particles and retain cell debris. The filter unit was
connected to a Masterflex pump, which accomplished a consistent flow rate of 40 ml/min.

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A. MCS Chromatography Purification Step

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The filtered virus suspension was then subjected to MCS chromatography. The MCS column was prepared as follows. 27 ml slurry led to 14 ml sedimentated resin which was packed using a Götec Superformance Column (diameter 1.0 cm, height 15.7 cm, volume 12.33 ml). 1% of the column volume of a 1% acetone solution was injected to the column and the column was run with a flow of 100 cm/h. The HETP, N and A_s values were then calculated as HETP: 0,056 cm. N/m: 1790 and $A_s = 1.20$.

The amount of proteins in the purified solution after the MCS chromatography step were assessed with a bicinchoninic acid (BCA) method (Interchim) (see, e.g., http://www.piercenet.com/files/bca.pdf) and electrophoresis.

SDS-PAGE was done in accordance to Laemmli, *Nature* (1970) 227:680-685. Samples for SDS-PAGE were diluted to a protein concentration of 77 μ g/ml. Different protein concentrations were loaded depending on the gel types used (10/12/15 Wells, Novex/Invitrogen):

Number of Wells	Protein Concentration in the Dilution	Load	Protein/Well
10 Wells	77 μg/ml	· 20 μl	1 μg
12 Wells	77 μg/ml	15-20 μ1	0.75 - 1 μg
15 Wells	77 μg/ml	10 μl	0.5 μg

Samples for use in a reducing SDS-PAGE were prepared as follows:

	26 μl sample or diluted sample
	+ 10 μl NuPage Sample Buffer (4x) SDS NP0003
	+ 4 μl TCEP Bondbreaker Solution 77720
×	(1:2 in MilliQ water)
Final Volume:	40 μl

The samples were heated for 10 minutes at 70°C or left at room temperature for 1 hour (leaving the samples at room temperature prevents the M protein of Corona Virus to coagulate/forming complexes), and then centrifuged for approximately one minute at 14,000 rpm in a table top centrifuge.

Markers for use on the gel were prepared as follows. Gel bands containing less than 1 μ g of proteins were easily visualised with the silver staining procedure using the Silver Staining Kit Protein, Plus One Staining Protocol (Pharmacia Biotech).

Western blotting was performed as follows. A semi-dry blotting technique was used to transfer the proteins from the SDS gel to a nitrocellulose membrane. The transfer was performed with a current of 0.8 mA/cm² for 1 hour. A rabbit polyclonal antibody against SARS virus was used to perform the immuno probing using the Western Breeze, Novex Chromogenic Western Blot Immunodetection Kit (Novex/Invitrogen).

The chromatogram of the inactivated SARS MCS capture step is depicted in FIGURE 27.

To estimate purity, MCS chromatography fractions were analysed by silver staining on NuPage

10% or 4-12% Bis-Tris-Gel (Novex) under reduced conditions, heated for 10 minutes at 70°C (Figure 28). The fractions were also analysed under the same conditions by western blot (Figure 29) to estimate purity, using PAK 11/03 SARS Cov 270603 neutralizing titer 1:512 (this antibody was used for this and subsequent western blots). Purity estimates are as follows:

Sample	Volume / ml	[Protein] / µg/ml	Total Protein / mg	Step Recovery Protein / %
Corona Harvest	100	2547.6	254.76	100
After Filtration = Load	100	2440.3	244.03	95.8
Flow Through	85	2321.4	197.32	77.5
Wash	49.32	468.5	23.11	9.1
Peak 1	12.12	252.7	3.062	1,2
Total Recovery	-	-	464.4	86.5

B. Density Gradient Ultracentrifugation Step

The eluted SARS virus fraction was then subjected to density gradient ultracentrifugation with a swinging bucket rotor to further purify the inactivated virus. 3 ml of MCS peak fraction were loaded onto a linear gradient (15-60% sucrose; 17 ml 15% and 17 ml 60% sucrose in gradient mixer). The separation was performed with a Beckman SW 28 rotor at 20,000 rpm for 2 hours.

The content of sucrose and protein in the linear density gradient ultracentrifugation fractions are depicted in the following table, the graph in figure 30 and the estimation of purity in figure 31:

Fraction	Fraction Size / ml	[Sucrose] / %	[Protein] / μg/ml
1	2	61	96.12
2	2	59.4	98.62
3	2	57.5	87.63
4	2	54.5	86.91
5	2	50.5	79.9
6	2	47.2	74.3
7	2	43.7	68.05
8	2	40.2	60.43
9	2	37.2	57.38
10	2	34	53.12
11	2	30	50.63
12	2	25.7	35.02
13	2	22.4	35.33
14	2	19.5	39.25
15	2	15.5	69.79
16	2	8.5	169.03
17	2	8.5	128.96

The protein concentration of fraction 11 (Figure 31 SDS-gel) was measured again against a standard curve prepared in 30% sucrose and lead to a protein concentration of 3.67 μ g/ml (0.05

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 μg on the gel). The M protein appears to be missing in this preparation possibily due to sample treatment procedure (heated samples).

There may be discrepancies in the protein concentration measurements in Table 2 due to sucrose interference with this assay.

5 EXAMPLE 3 -Production, Inactivation and Purification of Whole SARS Virus Using MCS Chromatography Resin Purification Followed by Density Gradient Ultracentrifugation

Inactivated SARS virus was prepared as described in Example above.

A. MCS Chromatography Purification Step

In this example, 200 ml of inactivated SARS virus harvest were subjected to MCS

10 chromatography. The chromatogram of the capture step of inactivated SARS virus purification with MCS is depicted in FIGURE 32, the protein recovery in the following table and the estimation of purity in FIGURE 33:

Sample	Volume / ml	[Protein] / μg/ml	Total Protein / mg	Step Recovery Protein / %
Corona Virus Harvest	200	2239.2	447.83	100
After Filtration = Load	200	2245.1	449.02	100.3
Flow Through	185	2126.3	393.37	87.8
Wash	49.32	450.1	22.2	5.0
Peak 1	4.43	1245.6	5.52	1.2
Total Recovery	-		421.08	93.7

B. Density Gradient Ultracentrifugation Step

3.5 ml of MCS peak fraction were then loaded onto a linear gradient (15-40% sucrose: 16 ml 15% and 16ml 40% sucrose in gradient mixer). The separation was performed with a Beckman SW 28 rotor at 20,000 rpm for 2 hours.

The content of sucrose and protein in the linear density gradient ultracentrifugation fractions are depicted in the following table and the graph in FIGURE 34:

Tube	Fraction Size / ml	[Sucrose] / %	[Protein] / μg/ml
1	2	40	45.86
2	2	39	45.68
3	2	37.5	44.14
4	2	35.5	37.82
5	2	33.5	34.48
6	2	31.5	31.76
7	2	30.5	29.49
8	2	28	30.87
9	2	25.5	31:7
10	2	23.5	26.74
11	2	21.75	23.58
12	2	20	35.33
13	2	18	96.38
14	2 .	14.5	523.79

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_15	2	8	941.97
16	2	8	696.7

Protein recovery is shown in the following table and the estimation of purity is shown in figure 35. Electron Micrograph pictures of density gradient fractions 8, 9 and 10 are shown in figure 36:

Step	Volume / ml	Protein / μg/ml	Total Protein / mg	Step Protein %
Load	3.5 ml	1245.6	4359.6	100
Bulk Protein Fractions	3.5 ml	720.8	4324.9	99.2
Viral Peak Fraction	8 ml	29.7	237.6	5.5
Total Recovery	-		4562.5	104.7

EXAMPLE 4 - Mouse Immunization with Inactivated SARS Virus

Mice were immunized subcutaneously on days 0, 14, and 28 with 5 µg BPL-inactivated SARS-CoV particles (BPL-SARS-CoV), either alone or together with Alum or MF59 as adjuvants. Serum was collected on days 0 (pre-immunization), 13 (post 1st immunization), 28 (post 2nd), and 35 (1 week post 3rd immunization). Neutralizing antibodies were assessed for blocking SARS-CoV infection of Vero cells in vitro. After 3 immunizations, neutralization titers were in the range 1:100-1:1000, which are levels similar to those present in the serum of SARS convalescent patients. As shown in the following table, the non-adjuvanted vaccine induced neutralizing antibody after the third immunization, and potency of this vaccine was enhanced significantly by including the adjuvants, with neutralizing antibody appearing after then 2nd immunization and overall titers increasing after then 3rd immunization:

_		Neutraliz	ation Titer	
Immunogen	pre	post 1st	post 2nd	post 3rd
BPL-SARS-CoV+MF59 (5 μg)	< 1:20	< 1:20	1:158	1:630
BPL-SARS-CoV+Alum (5 μg)	< 1:20	< 1:20	1:67	1:612
BPL-SARS-CoV (5 μg)	< 1:20	'< 1:20	< 1:20	1:71
PBS	< 1:20	< 1:20	< 1:20	< 1.20

EXAMPLE 5 - Balb/cMouse Immunization with Inactivated SARS Virus

A Balb/c mouse model for SARS infection has been developed (Subbarao et al. (2004), J.Virol., 78:3572-77. In this model, Balb/c mice are inoculated intranasally with 10⁴ TCID₅₀ of virus. At 48 hours post-inoculation, a 2-log increase in the TCID₅₀ virus titer can be detected in the lungs of infected mice. While virus replication is readily detected, the mice do not show any SARS disease symptoms and spontaneously clear the virus one week after inoculation. A decrease in virus titer in previously-immunized animals as compared to control animals demonstrates a protective effect of the vaccine being evaluated.

In this example, four Balb/c mice per group are immunized three times with 5 μ g BPL inactivated SARS-CoV (days 0, 14, 28) either alone or in combination with MF59 and

challenged with 10⁴ TCID₅₀ of SARS-CoV on day 43. Two days following virus challenge the mice are euthanized and SARS-CoV is quantified from nasal turbinates (NT) and lungs and the mean virus titer for each mouse is measured. Control groups received PBS alone, or an influenza virus vaccine (FLU) with or without MF59 adjuvant. Data were as follows (see also Figure 51), where four mice were tested per group and virus titers are expressed as log₁₀ TCID₅₀ per gram of tissue:

	Virus replication in lungs of challenged mice		Virus replication in nasal turbinates of challenged mice	
Immunogen	# infected/ Mean (± SE) # tested virus titer		# infected/ # tested	Mean (± SE) virus titer
PBS	4/4	6.3 ± 0.3	3/4	2.8 ± 0.35
MF-59 alone	4/4	6.1 ± 0.13	3/4	3.0 ± 0.38
FLU vaccine (5 μg)	4/4	6.3 ± 0.07	3/4	2.9 ± 0.36
FLU vaccine (5 μg) + MF-59	4/4	6.0 ± 0.19	4/4	3.0 ± 0.11
BPL-SARS-CoV (5 μg)	1/4	1.6 ± 0.13 *	0/4	Not detected **
BPL-SARS-CoV (5 μ g) + MF-59	0/4	Not detected *	0/4	Not detected **

Two-tailed Student's t-test, compared to PBS-immunized mice, showed: * P<0.00001 or ** P=0.025

As shown, virus could not be detected in the BPL-SARS-CoV immunized mice. The lower limit of detection of infectious virus in a 10% w/v suspension of lung homogenate was 1.5 log₁₀TCID₅₀/gm, and in a 5% w/v suspension of nasal turbinates the limit was 1.8 log₁₀TCID₅₀/gm. Viral titers in the immunized mammals were thus below these threshold values.

Thus the inactivated SARS-CoV vaccine was very efficient at preventing virus infection, as only one of eight mice immunized with the vaccine, either with or without MF59 adjuvant, was infected. Similar protection was not observed in control groups of PBS diluent, MF59 adjuvant, or influenza virus vaccine with or without adjuvant.

Neutralization titers of sera taken from the animals in the challenge study were assessed at two weeks post-1st, one week post-2nd, and one week post-3rd immunization. Mice immunized with the vaccine with MF59 adjuvant had already developed a neutralization titer of 1:71 after the 2nd immunization, which increased to 1:588 after the 3rd immunization, whereas mice receiving the unadjuvanted vaccine did not have any neutralizing activity post-2nd and a neutralization titer of 1:64 post-3rd immunization. Sera from mice in each of the control groups did not show any neutralization activity. These data clearly demonstrate not only the ability of the inactivated SARS-CoV vaccine to induce protective levels of SARS neutralizing antibodies, but also a beneficial effect of formulating the vaccine with adjuvant for elevated neutralization titers.

EXAMPLE 6 - Preparation of OMV comprising SARS viral antigens

E.coli were transfected with a plasmid of interest (encoding a SARS viral antigen). Single colonies harbouring the plasmid of interest were grown overnight at 37°C in 20 ml of LB/Amp

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(100 μ g/ml) liquid culture. Bacteria were diluted 1:30 in 1.0 L of fresh medium and grown at either 30°C or 37°C until the OD₅₅₀ reached 0.6-0.8. Expression of recombinant protein was induced with IPTG at a final concentration of 1.0 mM. After incubation for 3 hours, bacteria were harvested by centrifugation at 8 000 x g for 15 minutes at 4°C and resuspended in 20 ml of 20 mM Tris-HCl (pH 7.5) and complete protease inhibitors (Boehringer-MannheimTM). All subsequent procedures were performed at 4°C or on ice.

Cells were disrupted by sonication using a Branson Sonifier 450 and centrifuged at 5 000 x g for 20 min to sediment unbroken cells and inclusion bodies. The supernatant, containing membranes and cellular debris, was centrifuged at 50000g (Beckman Ti50, 29 000 rpm) for 75 min, washed with 20 mM Bis-tris propane (pH 6.5), 1.0 M NaCl, 10% (v/v) glycerol and sedimented again at 50000g for 75 minutes. The pellet was resuspended in 20mM Tris-HCl (pH 7.5), 2.0% (v/v) Sarkosyl, complete protease inhibitor (1.0 mM EDTA, final concentration) and incubated for 20 minutes to dissolve inner membrane. Cellular debris was pelleted by centrifugation at 5000g for 10 min and the supernatant centrifuged at 75000g for 75 minutes (Beckman Ti50, 33000 rpm). Outer membrane vesicles were washed with 20 mM Tris-HCl (pH 7.5) and centrifuged at 75 000 x g for 75 minutes or overnight. The OMV was finally resuspended in 500 µl of 20 mM Tris-HCl (pH 7.5), 10% v/v glycerol. Protein concentration was estimated by standard Bradford Assay (Bio-Rad), while protein concentration of inner membrane fraction was determined with the DC protein assay (Bio-Rad). Various fractions from the isolation procedure were assayed by SDS-PAGE.

EXAMPLE 7 - Immunogenicity, dose and route schedule for recombinant Spike protein in mice

The immunogenicity, route and dosing of the recombinant spike proteins of the invention in mice may be assessed using the below detailed protocol. Preferably, the administered antigen will elicit neutralizing antibody titers at least in the range of 1/100-1/1000. Increasing doses of antigen can be tested in the range from 5 to 20 μ g of recombinant Spike antigen alone or mixed with an equal volume of MF59-citrate, administered SC or IM to anesthetized mice in 100 μ l of inoculum. Groups of BALB/c mice, 6 per treatment are primed at day 0 and boosted at day 14 and 28.

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Group	Treatment	Dose/Route	Sampling interval	Number of mice
1-3	Rec-Spike protein	20, 10, 5 μg/SC	7, 21, 35, 42 d	6 per dose level
4-6	Rec-Spike protein	20, 10, 5 μg/SC	7	6 per dose level
7-9	Rec-Spike protein	20, 10, 5 μg/IM	7, 21, 35, 42 d	6 per dose level
10-12	Rec-Spike protein	20, 10, 5 μg/IM	7	6 per dose level
13-15	Rec-Spike - MF59	20, 10, 5 μg/SC	7, 21, 35, 42 d	6 per dose level
16-18	Rec-Spike - MF59	20, 10, 5 μg/SC	. 7	6 per dose level
19-21	Rec-Spike - MF59	20, 10, 5 μg/IM	7, 21, 35, 42 d	6 per dose level
22-24	Rec-Spike - MF59	20, 10, 5 μg/IM	7	6 per dose level
25	MF59	NA/SC	7, 21, 35, 42 d	6 + 6 (sac d 7 and 42)
27	MF59	NA/IM	7, 21, 35, 42 d	6 + 6 (sac d 7 and 42)
29	Saline	NA/SC -	7, 21, 35, 42 d	6 + 6 (sac d 7 and 42)
31	Saline	NA/IM	7, 21, 35,42 d	6 + 6 (sac d 7 and 42)

This protocol can also be used to assess the Th1/Th2 profile of the specific immune response elicited by the recombinant Spike protein. Neutralizing and Spike-specific antibody titers will be assessed at days 7, 21, and 35; IgG2a vs IgG1 isotype of the Spike-specific antibodies will be determined at days 21 and 35; in vitro proliferation of lymph node and splenic T cells against the recombinant Spike protein will be determined at days 7 and 42, respectively; IFN-γ and IL-4 production by splenic T cell against the recombinant Spike protein from SARS-CoV will be assessed at day 42. Peripheral blood will be collected at days 7, 21, 35; lymph nodes cells at day 7, and spleen cells at day 42. Neutralizing and Spike-specific antibody titers and isotypes will be determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively. Proliferation of lymph node and splenic cells will be determined by ³[H]—Thymidine uptake. Frequencies of splenic IFN-γ and IL-4 producing T lymphocytes, will be determined by ELISPOT and FACS.

EXAMPLE 8 -Immunogenicity, dosing and route schedule for Spike proteins in rabbits

The immunogenicity, route and dosing of the recombinant spike proteins of the invention in rabbits may be assessed using the below detailed protocol. Increasing doses can be tested in the range from 5 to 40 μ g of recombinant Spike antigen alone or mixed with an equal volume of MF59-citrate, administered SC or IM to anesthetized animals in 200 μ l of inoculum. Groups of New Zealand white female rabbits, 10 per treatment, will be immunized as shown in the table below. The animals will be primed at day 0 and boosted at days 14 and 28. Peripheral blood will be collected at days 7, 21, and 35. Neutralizing and Spike-specific antibody titers will be determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively.

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Group	Treatment	Dose/Route	Sampling interval	Number of rabbits
1-4	Full-length Spike protein	40, 20, 10, 5µg/SC	7, 21, 35 d	10 per dose level
5-8	Full-length Spike protein	40, 20, 10, 5µg/IM	7, 21, 35 d	10 per dose level
9-12	Truncated Spike protein	40, 20, 10, 5µg/SC	7, 21, 35 d	10 per dose level
13-16	Truncated Spike protein	40, 20, 10, 5µg/IM	7, 21, 35 d	10 per dose level
17-20	Full-length Spike protein - MF59	40, 20, 10, 5μg/SC	7, 21, 35 d	10 per dose level
21-24	Full-length Spike protein - MF59	40, 20, 10, 5µg/IM	7, 21, 35 d	10 per dose level
25-28	Truncated Spike protein - MF59	40, 20, 10, 5µg/SC	7, 21, 35 d	10 per dose level
29-32	Truncated Spike protein - MF59	40, 20, 10, 5µg/IM	7, 21, 35 d	10 per dose level
33	MF59	NA/SC	7, 21, 35 d	10
34	MF59	NA/IM	7, 21, 35 d	10
35	Saline	NA/SC	7, 21, 35 d	10
36	Saline	NA/IM	7, 21, 35 d	10

EXAMPLE 9 - Immunogenicity and dose schedule for recombinant Spike in ferrets

The immunogenicity and dosing of the recombinant spike proteins of the invention in ferrets may be assessed using the below detailed protocol. Three groups of ferrets, 6 for treatment, will be immunized with recombinant SARS-CoV Spike protein from CHO cell lines, alone or mixed with an equal volume of MF59-citrate, administered SC to anesthetized animals in 200µl of inoculum. The recombinant Spike protein vaccine will be tested at the dose eliciting the highest neutralizing antibody titers in mice at day 35 after the second boost. The animals will be primed at day 0 and boosted at day 14 and 28. Peripheral blood will be collected at days 7, 21, and 35. Neutralizing and Spike-specific antibodies titers will be determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively.

Groups	Treatment	Dose/Route	Sampling interval	Number of ferrets
1 & 2	Rec-Spike protein	Y μg or 2Y μg /SC	7, 21, 35 d	6
3 & 4	Rec-Spike protein + MF59	Y μg or 2Y μg/SC	7, 21, 35 d	6-
5	Saline	NA/SC	7, 21, 35 d	6

The 3 groups of ferrets, 6 animals per group, used for the immunogenicity studies above can then be used to assess efficacy of the recombinant Spike protein in protecting vaccinated animals from infection and/or disease. Anestethized animals will be challenged two weks after the last boost intratracheally with 10⁶ median tissue culture infectious dose unit (TCID₅₀) of the SARS-CoV Utah strain. Infection by SARS-CoV will be assessed by taking nasal, faringeal and rectal swabs from animals for 20 days after challenge as described (12). The presence of SARS-CoV in sample materials will be assessed by RT-PCR and infection assay of Vero cells.

Animals will be monitored for clinical signs of SARS disease by assessing sleeping time, temperature, respiratory symptoms, diarrhea, body weight and survival. Protection will be determined by the magnitude and duration of virus shedding and by duration and severity of disease symptoms and percentages of surviving animals.

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EXAMPLE 10: Expression of Spike protein for vaccination

The SARS-CoV Spike glycoprotein was expressed in both full-length and truncated forms, using the nSh and nShΔTC pCMVIII constructs described above, both with hexahistidine tags. The vector constructs were evaluated for expression 48 hr after transfection into 293 cells and COS7 cells. The full-length Spike protein (nSh) was detected by western blot only in cell lysate, but not in culture media (Figure 52).

The majority of SARS-CoV full-length Spike protein was expressed in transiently-transfected COS7 cells as a high molecular glycoprotein which ran at 540 kDa in non-reducing gels (Figure 53). The gp540 is heat labile as indicated by the complete dissociation into monomeric forms (gp170 & gp180) by boiling, but it was resistant to DTT treatment. These data suggest that the recombinant Spike protein is noncovalently associated into a homotrimer (gp540). The presence of Spike protein in homotrimeric association also was confirmed in inactivated, purified SARS-CoV virion particles. Analysis of virion proteins by western blot under the same condition used for the characterization of recombinant Spike protein generated essentially identical results (Figure 54).

EXAMPLE 11: Spike protein processing

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In order to characterize Spike protein processing, BHK-21 cells were infected with alphavirus replicon particles expressing the SARS-CoV full-length Spike. At 6 hoursr post-infection with an MOI of 5, infected cells were labeled for 1 hr with L-[35S]methionine/cysteine and chased for up to 4 hours. The [35S]-labeled spike protein was immunoprecipitated by anti-SARS rabbit serum and digested with Endo-H. Both digested and undigested proteins were analyzed by SDS-PAGE (4% polyacrylamide). As shown in Figure 55, the full-length spike protein is synthesized as an Endo-H sensitive high-mannose glycoprotein (gp170, an ER form) that undergoes modification to an Endo-H resistant glycoprotein with complex oligosaccharides (gp180, a Golgi form). The conversion of gp170 into the gp180 form takes place within 2 hours (Figure 56).

EXAMPLE 12: High-level protein expression

To develop a system for rapid expression of protein antigens, DNA transfection of 293 (human embryonic kidney) cells was used, to obtain milligram quantities of recombinant antigen. The most common method for culturing and transfecting 293 cells is in static or monolayer cultures. These procedures were modified by performing large-scale transfection of 293 cells in suspension and expanding the transfected cells in suspension culture for production of secreted or intracellular proteins. Several initial experiments were performed at the 100-milliliter scale cultures to determine optimum conditions, such as number of cells, type of

transfecting reagent (FuGENE 6, Lipitoid or RO-1538) and the ratio of DNA to transfection reagent. Based upon pilot experiments, FuGENE 6 was the best transfecting reagent.

The kinetics of gene expression was compared to other viral envelope glycoproteins, and the data suggest that stable protein expression peaks around 72 to 96 hours post-transfection, depending upon the gene of interest, and then significantly decreases thereafter. Thus, using the optimum conditions, the transfection process was scaled from 100 ml to 4 liters. The 4 liter culture can be used for rapidly producing 2-10 milligrams of protein antigens. To facilitate antigen purification and also maximize the yield and recovery of the purified protein, transfection conditions were optimized by using serum-free medium.

Bulk transfection procedure has been used for the expression of truncated and full-length Spike antigens. The kinetics of expression for truncated form of the spike protein is presented in Figure 56A. Expression of the truncated form of Spike protein peaked around 48 hrs and was stable until 72 hrs, therefore the cultures were harvested at 72 hrs post transfection.

Collected media were concentrated 20X and used for purification of truncated Spike protein by a very simple purification strategy where the truncated form of the spike was captured on GNA lectin followed by DEAE and ceramic hydroxyapatite column chromatography. The purified protein was analyzed on SDS-PAGE by silver stain (Figure 56B) and also by western blot (Figure 56C). Early efforts were able to purify the truncated form of the spike protein with >95% purity and approximately 50% recovery. The molecular mass of the truncated form of the Spike protein is approximately 170-180 kDa.

Full-length Spike protein was expressed in 293 cells using the bulk transfection strategy. The expression data suggest that, like the truncated form, expression peaked around 48 hrs post-transfection and remained stable until 72 hrs. However, contrary to the truncated form and as expected, full-length protein is not secreted, but rather is retained within the cells, as shown by the absence of any signal in western blots of cell culture supernatants. The full-length form of the protein was purified from Triton X-100 detergent-extracted cells. Full-length Spike protein was then captured on GNA lectin, followed by hydroxyapatite and SP chromatography. The calculated molecular mass of full-length spike protein is approximately 600 kDa, which is close to the theoretical mass for the trimer.

30 EXAMPLE 13: SARS virus seed cultures

A SARS-CoV reference seed virus propagated only in certified Vero cells will be used for the generation of the Master and Working Virus Seeds under GMP. A clinical specimen from the respiratory tract of a patient infected by the SARS-CoV is inoculated onto documented VERO cells, with certified culture media. Culture media containing the virus are harvested at 4 days post-infection and designated Passage 1 (P1). A second round of virus propagation is again

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performed in certified VERO cells with certified media, by inoculation of 1 ml per T-75 flask of 100 times diluted P1 virus. Culture supernatant was harvested at 3 days post-infection and stored at -80°C as a P2 reference stock virus, without plaque purification.

Cell banks of Vero cells for further production of SARS-CoV are prepared from specific cell subsets that have not been used since the emergence of transmissible spongiform enephalopathies (e.g. since 1980). A research cell bank of these cells has been prepared using specified New Zealand-origin fetal bovine serum. From this research cell bank, a Master Cell Bank (MCB) is made under GMP conditions and using only specified and well-controlled media and supplements. The cell bank will is tested for absence of adventitious agents according to applicable US, EU, and international guidelines (see Points To Consider "Characterization of cell lines used to produce biologicals", FDA/CBER 7/1993; ICH Q5D Draft 6 "Cell substrates", Oct.23, 1996; CPMP/ICH/294/95 "Note for Guidance on Quality of Biotechnological Products: Derivation and Characterisation of Cell Substrates Used for Production of Biotechnological/Biological Products (Step 4, 16. July 97); WHO final draft "Requirements for use of animal cells as in vitro substrates for the production of biologicals" 7.3.1997).

Tumorigenicity and identity testing is also required for this cell bank.

The reference virus is plaque-purified and expanded in certified Vero cells in the absence of FCS in order to generate Master and Working Seeds. Another option to help ensure purity and facilitate the assessment of safety of the Master Seed is to subject the SARS-CoV to pelleting and resuspension in PBS. The virus suspension is made up to 60% (w/w) sucrose with crystalline sucrose, transferred to a centrifuge tube and overlayed with 50, 40, 30, and 20% (w/w) sucrose solutions in PBS. The gradient is centrifuged for 72h and then fractionated. The virus-containing fraction is diluted and the virions re-pelleted by ultracentrifugation. RNA from the virus pellet is isolated and transfected into certified Vero cells whereby the "infectious" positive-strand RNA will lead to the production of infectious virus, which can be plaque-purified and expanded to generate alternative Master and Working Seeds from purified virus RNA.

Viral seeds are tested for the absence of adventitious agents (see e.g. 21 CFR Revised as of April 1, 1994, § 630.35 Test for safety) and for identity, using a highly-specific neutralizing antiserum prepared from an independent source. Safety testing of viral seeds for vaccine purposes is done routinely by service laboratories. Broad-spectrum PCR testing can be used as an addition and/or alternative for testing.

EXAMPLE 14: Scale-up of virus production and inactivation

A protocol for the production, inactivation, and purification of inactivated SARS-CoV with sufficient structural integrity to elicit protective neutralizing antibody responses in animal models involves: Vero cells are infected with virus at an M.O.I. of 0.01 in the absence of FCS

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and antibiotics; culture medium is collected, cleared by centrifugation, and inactivated with BPL, followed by confirmatory testing for complete inactivation; the inactivated material is filtered, subjected to MCS-column purification, and further purified by sucrose gradient centrifugation.

Several modifications and improvements can be developed when adapting this basic protocol to a larger scale for commercial use. Firstly, the cell culture and infection process can be adapted to roller bottles, as an intermediate step to allow rapid production for preliminary trials within existing BSL 3+ facilities. Full commercial production will typically use a fermentation process in a closed system, but a roller bottle system can be achieved more rapidly. The roller bottles do offer a true suspension culture system for Vero cells, which gives various technical and safety advantages over microcarrier cultures. Suspension cultures can be grown to any desired fermentation scale without interfering with the closed system between cell passages, as no trypsinization is required.

To scale up the infection process in roller bottles to 30-50 liters per batch, the optimum M.O.I. and harvesting periods for selected media and culture conditions should first be determined. For the larger scale, methods for harvesting and handling larger volumes of highly infectious material safely should be used, and so cell separation via centrifugation should be replaced by a method such as filtration through single-use filter cartridges.

The MCS-chromatography and the gradient purification steps described above can readily be scaled to a batch volume of up to 50 liters. For larger volumes, however, and for increased purity, ultrafiltration and sterile filtration steps will be used. Nuclease treatment to remove host cell DNA will also be included.

EXAMPLE 15: Large scale analytical methods

Analytical methods for the SARS coronavirus include virus titration methods, immunological and physico-chemical methods to quantitate and characterize the purified antigen (ELISA, PAGE, western blots using specific antisera against purified whole virus, etc.). Other analytical tests include: fast yield testing via asymmetric field flow separation and laser particle detection and counting; Western blot using specific antisera against individual viral proteins; and tests for residual host cell DNA.

Residual DNA testing is generally done by hybridization e.g. using a limit test. Such testing is performed according to methods already established and validated for other cell lines. As an alternative, the ThresholdTM method may be used.

For producing specific antibodies, recombinant protein expression of all the ORFs from the structural and non-structural gene regions of the SARS-CoV is used. The ORFs can be cloned and expressed in *E.coli* and, if necessary, also in eukaryotic vectors such as baculovirus. This can provide sufficient amounts of purified soluble protein to immunize mice and rabbits to produce

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polyclonal and monoclonal antibodies against SARS proteins and to set up specific ELISA assays. Different expression vectors can be tested to maximize the yield of recombinant protein in a soluble form e.g. different vectors, one containing sequences coding for six N-terminal histidine residues and another containing a Glutathione-S-transferase protein fused to the C-terminus of the SARS protein. The recombinant proteins can be purified by single step column chromatography on either Nickel chelating Sepharose or Glutathione-Sepharose 4B resin. These procedures are very rapid and generally produce protein of 60-90% purity, which is suitable for raising specific antisera (Pizza et al. (2000) Science 287:1816-20). Five mice and two rabbits for each recombinant protein can be immunized SC with 20 and 50 µg recombinant protein, respectively, given in IFA as adjuvant, at day 0, 14 and 28. Sera are collected at day 7, 21 and 35 to assess specific titers before euthanasia of the animals for collection of blood and removal of spleens.

For the detection of impurities (e.g. Vero cell derived proteins) in the vaccine preparation, rabbit serum reactive against Vero-derived proteins can be used. Such antisera are obtained by immunizing rabbits with at least 10µg of Vero cell lysate with CFA/IFA. The sera can be verified for reactivity against Vero-derived proteins in western blots. For more specific antisera against specific relevant cell-derived proteins that tend to be co-purified with the virus, mockinfected cell culture harvest that have undergone the purification process can be prepared and used for immunizing rabbits.

Methods to determine neutralization titers of sera from immunized animals and humans can be developed, without the constraints of using infectious SARS-CoV in a BSL-3+ laboratory. One such strategy will be to use recombinant antigens, particularly Spike protein or Spike-derived epitopes, and to develop ELISA assays for measuring antibodies against the target protein. Suitable epitopes allow a correlation to be established between the ELISA values and virus neutralization assay values. This approach provides a faster and more efficient (higher-throughput) comparison of specific and protective antibody titers. This ELISA test is also the ideal tool to monitor specific antibodies in safety trials, where several hundred animal sera must be tested.

Another strategy is to combine structural elements from both the pathogenic SARS-CoV and the non-pathogenic coronavirus mouse hepatitis virus (MHV) to construct chimeric virus-like particles (VLPs) that can be labeled. The assay is based on fusion between octadecyl rhodamine (R18)-labeled VLPs and cells (Hoekstra et al. (1984) Biochemistry 23:5675-81). The method relies on the relief of fluorescence self-quenching of R18 incorporated into VLPs upon fusion with cellular membranes. Coronavirus VLPs have been shown to mimic native virions with respect to their appearance in the electron microscope (EM) and their biological activities. As they do not contain viral RNA, however, then they cannot cause a productive infection

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(Vennema et al. (1996) EMBO J 15:2020-2028). The VLP system can be used for the mouse hepatitis virus (MHV) strain A59 (MHV-A59))Godeke et al. (2000) J Virol 74:1566-15) containing a chimeric S protein. The protein chimera, consisting of the ectodomain of the SARS-CoV and the transmembrane and endodomain (64 C-terminal amino acid residues) from the MHV spike protein, can be co-expressed with the MHV M (membrane) and E (envelope) protein in OST-7 cells) Godeke et al.). VLPs secreted in the supernatant are harvested, purified and labeled with octadecyl rhodamine (R18) (Hoekstra et al). A constant amount of VLPs is incubated with a serial dilution of sera at 37°C for 1 hour in a 96-well plate. Subsequently, cells expressing the receptor for the SARS-CoV, the angiotensin-converting enzyme 2 (ACE2) (Li et al. (2003) Nature 426:450-54) is be added and the extent of fusion can be measured with a fluorescence spectrophotometer.

A final strategy to monitor the ability of sera to inhibit cell-cell fusion interactions between cells expressing the SARS-CoV S protein and a human cell line expressing the angiotensin-converting enzyme 2 (ACE2), a functional receptor for SARS-CoV (Li et al.). This reporter gene-based assay uses the fluorescent shift (green to blue) of the fluorogenic substrate CCF2/AM (AM=acetoxymethyl) upon cleavage by β-lactamase (Bla) as read-out for cell-cell fusion (Zlokarnik et al. (1998) Science 279:84-88). For this assay, a BHK-derived cell line, stably expressing Bla and the SARS-CoV S protein is generated. In addition, a human cell line expressing ACE2 on its surface is used. BHK cells, expressing the S protein on their surface and Bla in their cytosol are incubated with serial dilutions of the sera to be tested for 1h at 37°C. The cell line expressing the ACE2 is loaded with 1μM CCF2/AM for 1 h at 22°C, washed twice with PBS, and co-cultivated with the BHK cells. In case of cell-cell fusion, Bla cleaves the substrate, resulting in a green blue shift with excitation at 409 nm. Inhibition of fusion by sera thus provides a detectable change.

EXAMPLE 16: Stabilisation of inactivated SARS-CoV

Although the purified inactivated SARS-CoV vaccine is capable of inducing potent neutralizing antibody responses in animals, it is relatively instable and can benefit from formulation to increase stability for an acceptable period of time. Suitable formulation changes include the use of various buffer systems, pH ranges, stabilizing excipients (e.g. sugars and sugar alcohols, amino acids, etc.) etc.. Stability testing can be conducted in real-time at normal storage temperatures, or can be conducted in an accelerated manner by using elevated temperatures. Vaccine stability can thus be increased to approximately one year or longer. Lyophilized vaccine formulation can also be used to extend shelf-life, possibly with further additives for stability during lyophilisation.

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EXAMPLE 17: Dose and schedule optimization for inactivated virus

Animal models of SARS-CoV infection have been reported, including mice, ferrets and macaques. As mentioned in example 4 above, mice immunized with the BPL-SARS-CoV vaccine achieve neutralizing antibody titers in the range of 1:100 – 1:1000, similar to levels found in convalescent patients, and are 100% protected from infection with a challenge virus. While the mouse challenge model is limited only to infection but not disease, ferrets and macaques are useful models of the human SARS disease. Two to four days after inoculation with SARS-CoV, both ferrets and macaques have been found to shed infectious SARS-CoV particles from the throat, nose and pharynx, as demonstrated by RT-PCR and/or virus isolation on Vero cells. At approximately the same time, the infected animals became lethargic, show respiratory distress and eventually die. Histologically, SARS-CoV infection in these animals associates with pulmonary lesions of different severity, similar to those found in biopsied lung tissue and autopsy material from SARS patients. With the availability of these models, preclinical studies with vaccines can be performed initially in mice for immunogenicity readouts, while efficacy of optimal doses and schedules can be assessed in the ferret and macaque models.

Initial studies in mice are used to determine the optimal dose and schedule required to elicit the highest levels of neutralizing antibody, with titers at least in the range of 1/100 – 1/1000. In parallel to the assessment of neutralizing activity, other features of the humoral immune response and cellular immune responses can be investigated. In particular sera from immunized mice can be assessed for the isotype (IgG1 vs. IgG2a) of the Spike-specific antibody response. Also, the frequencies of splenic CD4+ T cells producing IFN-γ and IL-4 in response to BPL-SARS-CoV particles will be assessed by ELISPOT and ELISA. These experiments can provide insight into the quality of the T cell response helping the priming of a protective antibody response.

Increasing vaccine doses can be tested (e.g. from 5 to 20 µg of BPL-SARS-CoV alone or mixed with an equal volume of MF59-citrate), administered SC to anesthetized mice in 100µl of inoculum. Groups of BALB/c mice, 10 per treatment, are immunized, with priming at day 0 and boosting at days 14 and 28. Secondary endpoints compare the kinetics of neutralizing vs. Spike-specific antibody titers and assess the Th1/Th2 profile of the specific immune response, and so neutralizing and Spike-specific antibody titers are assessed at days 7, 21, 35, and at 2, 3, 4, and 5 months after priming. The IgG2a and IgG1 titers of Spike-specific antibodies are determined at days 21, 35, and at 2, 3, 4, and 5 months after priming. Proliferation and IFN-γ and IL-4 production by splenic T cells against recombinant Spike protein from SARS-CoV are assessed at day 42, and at the end of the 5th month. Peripheral blood is collected at days 7, 21, 35, and at 2, 3, 4, and 5 months after priming. Spleen cells will be obtained at day 42 and at the end of the 5th month. Neutralizing and Spike-specific antibody titers and isotypes are determined by inhibition

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of infection of Vero cells and by ELISA, respectively. Proliferation of splenic cells is determined by ³[H]-thymidine uptake. Frequencies of splenic IFN-7 and IL-4 producing CD4⁺ T lymphocytes is determined by ELISPOT and FACS analysis.

Based on mouse results, the BPL-SARS-CoV vaccine can be tested in ferrets for the induction of protective neutralizing antibody titers. Ferrets are immunized according to a similar schedule as the mice and at the dose that elicits the highest neutralizing antibody titers in mice at day 35 after the second boost. Three groups of ferrets, 6 per treatment, are immunized with BPL-SARS-CoV, alone or mixed with an equal volume of MF59-citrate, administered SC to anesthetized animals in 200ul of inoculum. The animals are primed at day 0 and boosted at days 14 and 28. Peripheral blood is collected at days 7, 21, and 35. Neutralizing and Spike-specific antibodies titers are determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively. Each group of ferrets is used to assess efficacy of the BPL-SARS-CoV in protecting vaccinated animals from infection and/or disease. Anesthetized animals are challenged intratracheally, two weeks after the last boost, with 106 median tissue culture infectious dose units (TCID50) of the SARS-CoV CDC strain. Infection by SARS-CoV can be assessed by taking nasal, pharyngeal and rectal swabs from animals for 20 days after challenge (Martina et al. supra). The presence of SARS-CoV in sample materials can be assessed by RT-PCR and infection assay of Vero cells. Animals can be monitored for clinical signs of SARS disease by assessing sleeping time, temperature, respiratory symptoms, diarrhea, body weight and survival. Protection can be determined by the magnitude and duration of virus shedding, by duration and severity of disease symptoms, and by percentage of surviving animals. The formulation eliciting the highest neutralizing antibody titers at day 35 can then be tested against a two-fold higher dose of BPL-SARS-CoV given in the same formulation in the same regimen.

Additional studies can evaluate immunogenicity and efficacy of the candidate vaccine in non-human primates. Three groups of adult cynomolgus macaques, 4 per treatment, are immunized with BPL-SARS-CoV, alone or mixed with an equal volume of MF59-citrate, administered SC to anesthetized animals in 500μl of inoculum. The BPL-SARS-CoV vaccine can be tested at the dose eliciting the highest neutralizing antibody titers in ferrets at day 35 after the second boost. The animals are primed at day 0 and boosted at 3 and 6 weeks. Peripheral blood is collected at weeks 1, 4, and 7. A secondary endpoint is to assess the Th1/Th2 profile of the specific immune response. Neutralizing and Spike-specific antibody titers and frequencies of peripheral blood CD4+ T cells producing IFN-γ and IL-4 in response to the recombinant SARS-CoV Spike protein is thus assessed at weeks 1, 4, and 7. Neutralizing and Spike-specific antibody titers can be determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively. Intracellular cytokine staining and FACS analysis will be used to quantify IFN-γ- and IL-4-producing CD4⁺T cells. The macaques can also be used to assess efficacy of

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the BPL-SARS-CoV in protecting vaccinated animals from infection and/or disease. Anesthetized macaques can be challenged two weeks after the last boost with 10⁶ median tissue culture infectious dose unit (TCID₅₀) of the SARS-CoV CDC strain in a 5 ml volume. A few drops of the virus can also be administered on each of the conjunctiva, 0.5 ml in the nose and the remainder in the trachea. Infection by SARS-CoV can be assessed by taking nasal, pharyngeal, and rectal swabs, and feces from animals for 20 days after challenge (Fouchier et al. (20030 Nature 423:240). The presence of SARS-CoV in sample materials can be assessed by RT-PCR and infection assay of Vero cells. Animals can also be monitored for clinical signs of SARS disease by assessing sleeping time, temperature, respiratory symptoms, diarrhea, body weight and survival. Protection can be determined by the magnitude and duration of virus shedding, by duration and severity of disease symptoms, and by percentage of surviving animals.

Mice

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Group	Treatment	Dose/Route	Sampling interval	Number of mice
1-3	BPL-SARS-CoV	20, 10, 5 μg/SC	7, 21, 35 d; 2, 3, 4, 5 m;	10 per dose level
4-6	BPL-SARS-CoV	20, 10, 5 μg/SC	42 d	10 per dose level
7-9	BPL-SARS-CoV MF59	20, 10, 5 μg/SC	7, 21, 35 d; 2, 3, 4, 5 m;	10 per dose level
10-12	BPL-SARS-CoV MF59	20, 10, 5 μg/SC	42 d	10 per dose level
13	MF59	NA/SC	7, 21, 35 d; 2, 3, 4, 5 m;	10 + 10 (sacrificed at 42 d and end 5 m)
14	Saline	NA/SC	7, 21, 35 d; 2, 3, 4, 5 m;	10 + 10 (sacrificed at 42 d and end 5 m)

Ferrets

Group	Treatment	Route	Sampling interval	No. of ferrets
1	BPL-SARS-CoV -	SC	7, 21, 35 d	6
2	BPL-SARS-CoV-MF59	SC	7, 21, 35 d	6
3	Saline	SC	7, 21, 35 d	6

Macagues

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Group	Treatment	Route	Sampling interval	No. of macaques
1	BPL-SARS-CoV	SC	1,4, 7 w	4
2	BPL-SARS-CoV - MF59	SC	1,4, 7 w	4
3	Saline	SC	1,4, 7 w	4

EXAMPLE 18: Human T cell responses

As a prelude to initiation of clinical studies in humans, the reactivity of peripheral blood T lymphocytes from healthy donors with different HLA haplotypes can be assessed using the *in vitro* priming technique (Abrignani *et al.* (1990) *Proc Natl Acad Sci U S A* 87:6136-40). The aim of this study is to have a first indication of the immune-dominant T cell epitopes in SARS-CoV

proteins. Briefly PBMCs from 20 healthy donors with different HLA haplotypes will be cultured in medium containing 5% autologous serum, in the presence of different concentration of SARS-BPL-CoV particles in the range from 0.5 to 20 μg/ml. The expression of activation markers will be assessed after 24 and 48 hours. Frequencies of IFN-γ- and IL-4- producing T lymphocytes will be assessed after 12h and after 15 days in culture, in the presence of 100 U/ml recombinant human IL-2. Activated and cytokines producing CD4 T lymphocytes will be sorted and eventually cloned as single cells using FACS technologies. The CD4+ T cell repertoire from human subjects with different HLA will be assessed by proliferation assays of the CD4+ T cell lines and clones against autologous EBV-transformed cell lines loaded with 15-mer overlapping peptides from the most relevant structural and non structural protein of the SARS-CoV.

When moving to actual human trials, safety and immune responses will be evaluated in healthy adults following intramuscular immunization with escalating doses of the BPL-inactivated SARS-CoV vaccine, with MF59 adjuvant being included or omitted depending on preclinical data. Three/four immunizations will be given at 0, 1, 6 months in the first cohort, and at 0, 1, 2, 6 months and 0, 2, 6 weeks in the second and third cohorts respectively. The trial will be observer blind and placebo controlled. Subjects will be randomized into each dose level. Immune response parameters to be measured will include serum neutralizing antibodies, ELISA antibodies and peripheral blood IFN-gamma-producing CD4+ T cells by intracellular cytokine staining.

Group	Antigen dose (µg)	Administration schedule	No. treated subjects	No. subjects with placebo	Sampling interval
A1	10	0,1,6 months	18	6	0, 1, 2, 6, 7 mos
A2	20	0,1,6 months	18	6	0, 1, 2, 6, 7 mos
B1	10	0,1,2,6 months	18	12	0, 1, 2, 6, 7 mos
B2	20	0,1,2,6 months	18	12	0, 1, 2, 6, 7 mos
C1	10	0,2,6 weeks	18	12	0, 2, 6, 10, 30 wks
C2	20	0,2,6 weeks	18	12	0, 2, 6, 10, 30 wks

EXAMPLE 19: Selection of CHO cell lines for Spike protein expression

Methods for the derivation of Chinese Hamster Ovary (CHO) cell lines that stably express viral envelope glycoproteins that are conformationally intact, appropriately glycosylated and efficiently bind neutralizing antibodies are well established for HIV and HCV (Srivastava et al. (2002) J Virol 76:2835-47; Srivastava et al. (2003) J Virol 77:11244-259; Heile et al. (2000) J Virol 74:6885-92). The same techniques can be applied to SARS-CoV, to generate two different stable CHOK-1 cell lines producing either full-length or truncated SARS Spike proteins. The Spike proteins can be expressed using the constructs described herein, but without the hexa-His tags. These proteins can compared for their ability to produce neutralizing antibodies in immunized animals as well as for their expression levels in CHOK-1 cells.

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A pCMV3 vector expressing Spike can be used for the derivation of stable CHOK-1 cell lines, containing the CMV enhancer/promoter, ampicillin resistance, and a fused DHFR and attenuated neomycin gene for selection purposes. Stable cell lines can produced using the neomycin selection system in CHOK-1 cells. Clones can be sequenced to verify the integrity of the insert, and transient transfections can be performed using Trans-LT1 polyamine transfection reagent (PanVera Corp., Madison, WI) to assess the expression level and also the integrity of the expressed protein by ELISA and western blot analysis.

Initial CHO cells will be selected to be free from TSE/BSE contaminants and risks according to relevant regulatory standards. To construct cell lines, procedures involve transfection, primary screening with selective medium, followed by subcloning to assure purity of cell lines. Cell supernatants can be assayed using an antigen capture ELISA to quantify expression levels at all stages of selection and amplification. For full-length Spike expression, methanol fixed cells can be screened for internal expression by immunofluorescent staining using a rabbit anti-SARS antibody. Successive measurements at the T75-flask stage of expansion canbe employed to assure stability of expression levels. The molecular mass and integrity of the expressed proteins can be checked by PAGE both under native and reducing and denaturing conditions, followed by immunoprobing.

The pCMV3 vectors expressing SARS-CoV Spike proteins in either full-length or truncated forms can be introduced into CHOK-1 cells using the Trans-LT-1 reagent and nonselective media. 24-48 hours post-transfection, depending on cell density, cells are split at a 1:5 ratio and the medium can be changed to selective media containing neomycin at 500µg/ml. Any bovine serum used in these procedures will be from TSE-free sources that meet regulatory standards. Ten to fourteen days later, individual colonies can be picked and transferred to 96 well plates and cultured in complete non-selective medium. When approximately 80% of the wells are confluent. 24 hour supernatants can be screened by Spike capture ELISA. For initial expression of full length Spike protein, cells can be fixed with methanol and screened by immunofluorescent staining using a rabbit anti-SARS antibody. After low-expressing cell lines have been eliminated and there are fewer than 20-30 cell lines, capture ELISA and western blots can then be used to determine the expression level after cell lysis. A portion of each cell line can be pelleted, weighed and lysed in 1% Triton lysis buffer for determination of expression levels. Three to four clones producing the highest levels of spike protein in correct structure and conformation can be expanded to three-liter bioreactors and adapted to low serum suspension culture conditions for scale-up.

The antigen capture ELISA assay for the SARS spike protein can be performed using 96 well flat-bottom plates coated with 250ng per well of purified immunoglobulin obtained from rabbit sera that were immunized with inactivated SARS virus. Supernatant or lysate samples are

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added and incubated for 2 hours at 37°C. Bound antigen is reacted against pooled SARS*ve serum or high affinity monoclonal antibody either human or mouse against SARS spike protein and detected using appropriate species-specific peroxidase-conjugated second antibody. The plates are developed using TMB substrate (Pierce, Rockford, IL), read at a wavelength of 450nm, and the concentration of protein per ml sample is derived from a standard curve (OD vs. protein concentration) based on serial dilutions of a known concentration of recombinant spike protein.

The immunoprobing analysis will also be performed following the standard methods described by Srivastava et al. (2002) supra. Briefly, 10-20µl of the sample is analyzed on 4-20% SDS PAGE under non-reducing/denaturing conditions with mild heating. The proteins are then transferred onto nitrocellulose membranes and reacted against polyclonal anti-Spike rabbit serum, followed by anti-rabbit Ig conjugated to Alexa 688 (Molecular Probes, Oregon). The blots are scanned using an infrared imaging system.

The highest expressing candidate cell lines will be screened for Spike protein expression and stability in small-scale (3 liter) perfusion bioreactors. The candidate clones will be further evaluated for level of expression as well as integrity of expressed protein, and subsequently tested for expression stability in the absence of selection. The selected clones also will be tested for maintenance of the DNA sequence integrity of the integrated SARS spike protein gene. To quickly monitor the expression levels in small flasks and in the three liter evaluation cultures, a lectin-based process (Gluvanthus Nivalis lectin) has been developed to isolate SARS spike protein to a degree of purity that allows semi-quantitation and characterization of the protein in CHO supernatant. Full-length Spike protein will be obtained from Triton X-100 detergent extracted cells and then captured on GNA lectin, followed by hydroxyapatite and SP chromatograph. Eluted protein is then characterized by: (1) polyacrylamide gel electrophoresis (PAGE) and Coomassie staining, (2) immunoprobing with anti-SARS rabbit sera, (3) structural characterization using size exclusion chromatography (SEC), as well as mass spec analysis using MALDI-TOF.

Productivity from the CHO cell line expressing SARS spike protein should be at least 2 mg/L and for full-length Spike protein will be 3mg/100gm of cells, at steady-state cell density. Yield from one 45 day, 2.5-liter bioreactor will be ~1000 mg crude protein.

EXAMPLE 20: Purification of spike protein for human vaccines

To purify SARS spike protein for the purpose of producing GMP grade material for human use, the following basic process is used, with all steps being performed at 2-8°C: the starting material, concentrated CHO cell culture supernatant (20-30X) is thawed and filtered through a 0.45µm membrane; this material is heavily contaminated proteins from culture, as well as DNA;

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the first purification step is affinity chromatography using Gluvanthus Nivalis (GNA), a lectin that preferentially recognizes terminal mannose containing carbohydrates; glycosylated proteins, including SARS spike protein are captured and non-glycosylated proteins, as well as DNA, do not bind to this column; the GNA column is followed by two chromatographic steps operated in the flow through mode; the anion exchanger, DEAE, and ceramic hydroxyapatite (cHAP); DEAE binds some contaminating supernatant proteins and DNA, whereas cHAP binds any contaminating serum proteins; full-length Spike protein is purified from the cell pellet; the cells are lysed with Triton X-100 and full-length Spike protein is then captured on GNA lectin, followed by hydroxyapatite and SP chromatography.

The purified SARS spike can be further treated to remove adventitious viruses: viral inactivation at pH 3.5 for 1 hour; the sample is then concentrated and diafiltered into a buffer at pH 4 and finally captured the purified protein using SP resin; the spike protein binds to this resin and many viruses flow through.

The spike protein is eluted, concentrated and diafiltered into formulation buffer. This formulated bulk product is then filtered through a DV50 viral removal membrane followed by filtration through a $0.2 \mu m$ membrane. The formulated bulk is filled into suitable containers e.g. into 3.0 ml vials, in a class 100 laminar flow hood.

In process testing at each step of the purification includes protein concentration, endotoxin (LAL), bioburden, and recovery.

Prior to human administration, a test for potency will evaluate the specific ability of the vaccine in an *in vitro* or *in vivo* test to effect a given response. The *in vivo* immunogenicity will be determined by dosing groups of 10 mice with various doses of the protein antigen. Sera will be analyzed for the presence of IgG antibodies using an ELISA. The criterion for passing will be based upon the number of vaccine treated animals that are seropositive compared to a reference standard. Other tests include General Safety, sterility, purity, identity of the vaccine (using an ELISA specific for Spike protein), and quantity & protein concentration (UV spectrophotometric absorbance procedure based on the molar absorbance of the aromatic amino acids).

Stability testing will be performed on the bulk drug substance and on the final container product. Bulk product will be evaluated at temperatures of -60° C (recommended storage condition), $25 \pm 2^{\circ}$ C and $40 \pm 2^{\circ}$ C protected from light, at time points of 0, 3, 6, 9, 12 months. Final container product will be tested at temperatures of -60° C, and inverted at $5 \pm 3^{\circ}$ C, $25 \pm 2^{\circ}$ C, and $40 \pm 2^{\circ}$ C at time points of 0, 3, 6, 9, 12 months. Stability-indicating assays may include appearance, pH, protein content, SDS-PAGE, size exclusion HPLC, and container/closure integrity, performed on single samples of bulk and triplicate vials of final container material.

The protein purified in this way can be evaluated in mice, rabbits and ferrets as described in, and based on the results of, examples 4, 5, 8 and 9 above.

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Initial experiments will be performed in mice to determine optimal dose and schedule of the GMP Spike protein required to elicit the highest levels of neutralizing antibody, with titers at least in the range of 1/100 - 1/1000. Spike protein will be tested in the range from 5 to 40 µg. alone or mixed with an equal volume of MF59-citrate, to anesthetized mice in 100ul of inoculum. Groups of BALB/c mice, 10 per treatment, will be immunized. The animals will be primed at day 0 and boosted at days 14 and 28. Secondary endpoints will be to compare the kinetics of neutralizing vs. Spike-specific antibody titers and to assess the Th1/Th2 profile of the specific immune response. Neutralizing and Spike-specific antibody titers will be assessed at days 7, 21, and 35 and at 2, 3, 4, and 5 months after priming; the IgG2a and IgG1 titers of Spikespecific antibodies will be determined at days 21 and 35, and at 2, 3, 4, and 5 months after priming; proliferation and IFN-y and IL-4 production by splenic T cell against the recombinant Spike protein from SARS-CoV will be assessed at day 42 and at the end of the 5th month. Peripheral blood will be collected at days 7, 21, and 35 and at 2, 3, 4, and 5 months after priming; spleen cells at day 42 and at the end of the 5th month. Neutralizing and Spike-specific antibody titers and isotypes will be determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively. Proliferation of splenic cells will be determined by 3[H]thymidine uptake. Frequencies of splenic IFN-y and IL-4 producing CD4+ T lymphocytes, will be determined by ELISPOT and FACS analysis.

Next, the optimal dosing and schedule for recombinant Spike vaccine will be determined in ferrets. Based on the mouse results, the Spike vaccine eliciting the highest antibody neutralizing titers will be tested against a two-fold higher dose of recombinant Spike protein given in the same formulation. Three groups of ferrets, 6 per treatment, will be immunized SC under anesthesia with 200μ l of inoculum. The animals will be primed at day 0 and boosted at days 14 and 28. Peripheral blood will be collected at days 7, 21, and 35. Neutralizing and Spike-specific antibodies titers will be determined by inhibition of SARS-CoV infection of Vero cells and by ELISA, respectively. Similar to the previous ferret studies, each group of animals will be used to assess efficacy of the vaccine in protecting immunized animals from infection and/or disease.

Immunogenicity and efficacy of the candidate vaccine also will be evaluated in nonhuman primates. Three groups of adult cynomolgus macaques, 4 per treatment, will be immunized with recombinant SARS-CoV Spike protein, alone or mixed with an equal volume of MF59-citrate, administered SC to anesthetized animals in 500 μ l of inoculum. The Spike protein vaccine will be tested at the dose eliciting the highest neutralizing antibody titers in ferrets at day 35. The animals will be primed at day 0 and boosted at 3 and 6 weeks. Peripheral blood will be collected at weeks 1, 4, and 7. A secondary endpoint will be to assess the Th1/Th2 profile of the specific immune response, as described above (neutralizing and Spike-specific antibody titers,

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frequencies of peripheral blood CD4+ T cells producing IFN- γ and IL-4 in response to the recombinant Spike protein, assessed at at weeks 1, 4, and 7).

Finally, human phase I, placebo-controlled, dose-escalation, safety/ immunogenicity trials will be performed for the IM recombinant SARS vaccine with MF59 adjuvant. The trial will evaluate safety and immune responses in healthy adults following immunization with escalating doses of SARS recombinant vaccine with MF59 adjuvant, administered intramuscularly. Three/four immunizations will be given at 0, 1, 6 months. The trial will be observer blind and placebo controlled. Subjects will be randomized into each dose level. Immune response parameters to be measured include serum neutralizing antibodies, ELISA antibodies and peripheral blood IFN-γ-producing CD4+ T cells by intracellular cytokine staining:

Group	Vaccine Antigen dose (µg)	Administration schedule	No. of treated subjects	No. of subjects with placebo (MF59)	Sampling interval
A1	50	0,1,6 months	18	6	0, 1, 2, 6, 7 months
A2	100	0,1,6 months	18	6	0, 1, 2, 6, 7 months

EXAMPLE 21: Comparison of inactivated virus and purified Spike protein

Immunogenicity and efficacy of the inactivated virus vaccine and the purified Spike protein can be compared in non-human primates. Three groups of adult cynomolgus macaques, 4 for treatment, will be immunized with recombinant SARS-CoV Spike protein from CHO cell lines or with BPL-SARS-COV, given in the dose and formulation eliciting the highest neutralizing antibody titers in previous immunogenicity challenge experiments, administered SC to anesthetized animals in 500μ l of inoculum. The animals will be primed at day 0 and boosted at 3 and 6 weeks. Peripheral blood will be collected at weeks 1, 4, 7. A secondary endpoint will be to assess the Th1/Th2 profile of the specific immune response, as described above.

-	Group	Treatment	Dose/Route	Sampling interval	No. of macaques
1	1	Rec-Spike protein	Y μg /SC	1,4, 7 w	4
		+ or - MF59		,	
	2	BPL-SARS-CoV	Y μg/SC	1,4, 7 w	4
		+ or - MF59			
	3	Saline	NA/SC	1,4, 7 w	4

EXAMPLE 22: Expression in yeast

Yeast is a useful and inexpensive eukaryotic expression system. Yeast-expressed proteins are used in recombinant hepatitis B virus vaccines, and recombinant SARS antigens may also be expressed in yeast for vaccine purposes. Yeast-expression is also convenient for the production of antigens for preparing monoclonal and polyclonal antitobodies, or for use in serological assays.

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The nucleocapsid protein (N) and two different versions of the spike glycoprotein (S) from SARS coronavirus FRA strain (AY310120) were cloned for expression in S.cerevisiae:

SARS N: aa 1 – 422 (coordinates 28120-29388 of AY310120 strain) – Fig.65 SARS spike: aa 14 – 1195 (transmembrane domain and cytoplasmic tail deleted) – Fig.66 SARS spike: aa 14 – 662 (S1 domain)

To make the S1 construct, a XhoI-NotI fragment of approximately 3733bp encoding the full-length spike glycoprotein was the starting point. PCR was used to amplify the full-length gene in two pieces: XbaI-BlnI of 2440bp and BlnI-SalI of 1306bp. These fragments were subcloned into commercial vectors (Novagen): pT7Blue2 XbaI-BlnI (5' end of spike glycoprotein) and pT7Blue2 BlnI-SalI (3' end of spike glycoprotein; Figure 58), respectively. The following primers were used in the subsequent PCR reactions: Spk-1 (5') SEQ ID NO: 9785; Spk-2 (5') SEQ ID NO: 9786; Spk-3 (5') SEQ ID NO: 9787, Spk-4 (5') SEQ ID NO: 9788.

E. coli HB101 competent cells were transformed with the PCR ligation product and plated on Luria agar plates, containing 100µg/ml ampicillin. The desired clones were identified using miniscreen DNA analysis. After sequence verification and plasmid amplification of the desired subclones, it was desirable to eliminate the internal SalI site present in the XbaI-BlnI portion of the spike sequence in order to facilitate future cloning into the yeast expression vector (BamHI-SalI). Therefore, we prepared a CelII-MfeI vector from the pT7Blue2 XbaI-BlnI (5' end Spike) subclone to eliminated a 143bp sequence containing the SalI site. Kinased oligos DS1-6 (SEQ ID NOS: 9789-9794) were then ligated into the CelII-MfeI vector to replace the 143bp that were removed to mutate the SalI site (no aa changes), creating pT7Blue2.XbaI-BlnIAsal.

The 5' XbaI-BlnI (from pT7Blue2.XbaI-BlnI \(\Delta Sal \)) and the 3' BlnI-SalI (from pT7Blue2 BlnI-SalI) spike glycoprotein inserts were gel-purified and ligated them into the p893-1 XbaI-SalI vector (a vector derived from pLitmus 38 (New England Biolabs) with the alpha-factor leader sequence cloned into the BamHI-SalI sites of the MCS). The resulting full-length SARS Spike coding sequence was named p893-1.SARS Spike 1255 #9 (Figure 58).

E.coli HB101 competent cells were transformed with the oligo replacement ligation product and plated on Luria agar plates, containing 100μg/ml ampicillin. The desired clones were identified using miniscreen DNA analysis. After sequence verification of the positive clones, pT7Blue2 Xba-Bln ΔSal was chosen for use as a template for PCR reactions to amplify the Spike S1 1967 bp Xba-Sal fragment. The fragment was then subcloned into the p893-1 Xba-Sal vector, sequence verified, and named it p893-1. Spike S1 #11 (Figure 59).

In order to clone into the *S.cerevisiae* expression vector, pBS24.1, the 5' end of the S1 sequence had to be modified from XbaI to HindIII to allow ligation with the 3' HindIII end of the ADH2/GAPDH BamHI-HindIII promoter fragment. From pT7Blue2 Xba-BlnΔSal (described above) an AgeI-SalI 1943bp fragment was gel-purified. This fragment was ligated along with a

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synthetic pair of HindIII-Agel 30bp kinased oligos (S1-1+S1-2 creating the necessary 5' HindIII site) into the pSP72 HindIII-SalI commercial subcloning vector (named pSP72.SARS Spike S1 #2; Figure 59). S1-1 had SEQ ID NO: 9795 and S1-2 has SEQ ID NO: 9796.

After sequence verification of the positive clone from miniscreen DNA analysis, the HindIII-SalI fragment was gel purified. The 1365 bp BamHI-HindIII ADH2/GAPDH promoter fragment was ligated along with the 1973 bp HindIII-SalI S1 fragment into the pBS24.1 BamHI-SalI vector creating the genetically engineered pd.SARS Spike S1 #2 expression plasmid (Figure 60).

S.cerevisiae strain AD3 was transformed with pd.SARS Spike S1 #2 and single transformants were checked for expression after depletion of glucose in the medium. The recombinant protein was expressed at high levels in yeast, as detected by Coomassie blue staining. In particular, yeast cells were transformed with the SARS S1 expression plasmid using the Invitrogen S.c. EasyCompTM Transformation Kit. Expression in shown in Figure 57.

To express Spike 1195 protein, which does not contain the trans-membrane (TM) region or cytoplasmic tail that are present in the full-length SARS construct, the following series of genetic manipulations was performed:

From pT7Blue2 BlnI-SalI #11 (described above) a BlnI-DraI 1056bp fragment was gel purified. This fragment was ligated with a synthetic pair of 68bp DraI-SalI kinased oligos (DRS1+2; SEQ ID NOS: 9797 & 9798) into a pT7Blue2 BlnI-SalI vector (Figure 61). E.coli HB101 competent cells were transformed with the oligo replacement ligation product and plated on Luria agar plates, containing 100µg/ml ampicillin. The desired clones were identified using miniscreen DNA analysis. After sequence confirmation the clone was named pT7Blue2 BlnI-Sal Spike 1195 #7. The 1126bp BlnI-SalI fragment encoding the 3' end of the Spike 1195 was gel purified (Fig.61).

In order to generate the XbaI-SalI Spike 1195 fragment, the 3109bp XbaI-PciI fragment was isolated from the p893-1.SARS Spike 1255 #9 (described above) and a 457bp PciI-SalI fragment from pT7Blue2.SARS Spike 1195 #7 (described above). The two fragments were cloned into the p893-1 XbaI-SalI vector, creating the p893-1.SARS Spike 1195 #34 plasmid (Figure 62).

To clone SARS Spike 1195 into the pBS24.1 Saccharomyces cerevisiae expression vector, it was necessary to modify the 5' end of the SARS Spike 1195 from XbaI to HindIII, as done for the Spike S1 expression clone described above. To begin, the 2416bp AgeI-BlnI fragment was isolated from p893-1.SARS Spike 1195 #34. This fragment was ligated with the synthetic HindIII-AgeI 30bp oligos (described above to generate the S1 protein for expression in S.cerevisiae) into the pT7Blue2 HindIII-BlnI vector. E. coli HB101 competent cells were transformed with the oligo replacement ligation product and plated on Luria agar plates,

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containing 100µg/ml ampicillin. The desired clones were identified using miniscreen DNA analysis. After sequence verification of the positive clone and plasmid amplification of pT7Blue2.SARS 1195 5' HindIII-BlnI #10 (Figure 63), we isolated a 402bp HindIII-NcoI fragment and the 2044bp NcoI-BlnI fragment (Figure 63). It was necessary for the HindIII-BlnI isolation to be done in two steps to avoid cloning issues related to the internal HindIII site located at nucleotide number 1319 of the spike 1195 protein.

To assemble the BamHI-SalI-expression cassette of Spike 1195 into the pBS24.1 vector E.coli HB101 competent cells were transformed with the the BamHI-HindIII (ADH2/GAPDH promoter), HindIII-NcoI 402bp fragment, NcoI-BlnI 2044bp and the BlnI-SalI 1126bp fragments into the pBS24.1 BamHI-SalI vector. The samples were plated on Luria agar plates, containing $100\mu g/ml$ ampicillin. The desired clone was identified using miniscreen DNA analysis, thus creating the genetically engineered pd.SARS Spike 1195 #10 (Figure 64).

S.cerevisiae strain AD3 was transformed with pd.SARS Spike 1195 #10 and single transformants were checked for expression after depletion of glucose in the medium. The recombinant protein was detected by Coomassie blue staining. In particular, yeast cells were transformed with the SARS 1195 expression plasmid using the Invitrogen S.c. EasyCompTTM Transformation Kit.

EXAMPLE 23: Expression in mammalian cell lines

cDNA fragments containing the S protein ORF of 1255 amino acids were amplified by RT-PCR from SARS viral RNA (Frankfurt isolate) grown in Vero cells. The amplified PCR fragments were cloned into pBlueScript vector, sequenced, and consensus spike sequence was assembled to create a full-length SARS spike clone, pBSnSh. *In vitro* transcription of pBSnSh followed by translation in a rabbit reticulocyte lysate resulted in the production of single polypeptide with an estimated molecular mass of ~140 kDa.

The insert of this plasmid was recloned via XhoI and Not I into a mammlian expression vector pCMVIII (Srivastava et al. (2003) J. Virol. 77:11244-11259) to create a construct, nSh (Fig. 74A). A PCR fragment containing a spike protein of 1195 amino acid, which was deleted for transmembrane (TM) domain and cystein-rich cytoplasmic tail (Cy) was amplified and cloned pCMVIII vector to generate the contstruct nShΔTC (Figure.74B). Both constructs were tagged with six histidine residues at the C-terminus in order to aid in their characterization. The Xho I/Not I fragment without a histidine tag also was subcloned into the alphavirus replicon vector backbone pVCRchim2.1 for use in the production of an alphavirus replicon particle chimera that expresses S protein. Production and characterization of the replication defective alphavirus vector particles was performed essentially as described previously (Perri et al. (2003)

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J. Virol. 77:10394-10403; Polo et al. (1999) PNAS USA. 96:4598-4603). The resultant alphavirus vector particles were named as VEE/SIN.

COS7 cells and BHK-21 cells were maintained in Dulbecco's modified Eagle's medium supplemented with 10% fetal bovine serum at 37°C and 5% CO₂ in air. COS7 cells were transfected with expression plasmids (nSh, nSh\DeltaTC) using a transfection kit (TransIt-COS, Mirus) following the manufacturer's protocol. The cells were washed once with ice-cold PBS and lysed with 1x Lysis buffer (20mM MOPS, 10mM NaCl, 1.5mM MgCl₂, and 1% Triton X-100) containing complete mini protease inhibitor (Roche). After a 30-min incubation on ice, the debris was cleared by centrifugation. The cleared lysate was either purified or used directly in western blotting.

To purify secreted spike proteins, medium from transfected cells was collected and subjected to centrifugation at 12,000 rpm for 10 min to remove cellular debris. The cleared medium was applied to a ConA-agarose column (Vector Lab). The column was washed extensively with 20mM sodium phosphate buffer, and then the bound proteins were eluted with 1M methyl α -D-mannopyranoside (MMP), 1M NaCl in 20mM sodium phosphate buffer. Column fractions containing SARS-CoV spike proteins were applied to MagneHis Protein purification system (Promega) following the protocol suggested by the manufacturer.

For western blot analysis, proteins were separated by 4-20% SDS-PAGE and then transferred electrophoretically to nitrocellulose membrane (Invitrogen). Membrane was blocked in blocking buffer (5% skim milk and 0.1% Tween 20 in PBS) and incubated with indicated antibody at room temperature for 1 hr, washed and probed with horseradish peroxidase (HRP)-conjugated secondary antibody (Biosource) followed by chemiluminescence (ECL system, Amersham) and exposed by X-ray films. The antibodies used were a mouse monoclonal antihistidine antibody (anti-His*tag Mab, Novagen), a rabbit polyclonlal antipeptide antibody against SARS-CoV spike proten (SmPab, Abgent), or rabbit anti-SARS sera (2BE) obtained by immunization of rabbits with purified SARS-CoV virion. The latter has a cell culture neutralizing titer of 1/2,500. Unless stated otherwise, antibody was used at 1/1,000 for antihistidine antibody and SmPab and 1/10,000 for anti-SARS rabbit sera.

Some spike proteins were treated with Peptide-N glycosidase F (PNGase F). Cell lysates were diluted in 0.5% SDS and 1% β -mercaptoethanol and denatured at 100°C for 10 min. After 2-fold dilution with 1% NP-40 in 50mM sodium phosphate (pH 7.5), the samples were treated with PNGase F (NEB) at 37°C for 1 hr. Enzyme-treated samples were analyzed by 4-12% SDS-PAGE in reducing condition. For a partial digestion with the PNGase, the cell lysates were diluted with 50mM sodium phosphate (pH 6.0) containing 0.75% Triton-X and treated with PNGase F (Calbiochem) at 37°C for 3 hr. Enzyme-treated samples were analyzed by 4-20% SDS-PAGE in nonreducing condition.

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Western blots of cells 48-hours after transfection are shown in Figure 75. The S protein was detected in cell lysates as a doublet with estimated molecular weight of ~170 ~180 kDa , when the lysate was boiled and analysed under reducing SDS-PAGE conditions (Fig. 75A, lane 3). This doublet appears to result from differential glycosylation of one polypeptide product since pre-treatment of the cell lysate with PNGase F reduced the doublet to a single species of ~140 kDa (Fig. 75A, lane 4). This is the expected size predicted from the aa sequence for a full-length, intact polypeptide product. This experiment indicates that the full length SARS-CoV S is expressed in mammalian cells as a single, uncleaved polypeptide, but in two differentially glycosylated forms, gp170 and gp180 respectively. Unlike the two S glycoforms encoded by the full-length sequence, none of which were secreted, the S Δ protein product was detected both in cell lysates (Fig. 75A, lane 5) as well as in the cell culture medium (Fig. 75B, lane3) as a single species of ~160 kDa.

In order to further characterize the intracellular processing of the S protein, and as described above, BHK21 cells were infected with defective alphavirus particles expressing the full-length S. At 6 hr post infection with a MOI of 5, infected cells were pulse labeled for 1 hr with L-[35S] methionine/cysteine and chased for 2 or 4 hours. The [35S]-labeled S protein was immuno-precipitated using the rabbit antiserum raised against inactivated, purified virus and then digested with Endo H. The Endo H treatment involved dilution with a sample buffer (50mM sodium phosphate, 0.1% SDS, 50 mM DTT, pH 6.0) and boiling for 5 min. After denaturation, the samples were further diluted with 0.75% Triton-X 100 and treated with endoglycosidase H (Endo H) following manufacturer's protocol (Calbiochem) for 3 hr at 37°C. Enzyme-treated samples were added with gel loading buffer containing 0.1% SDS and DTT and analyzed by 8% SDS-PAGE.

Both digested and undigested proteins were boiled in SDS and analysed by reducing SDS-PAGE (Figure 55). After a 1-hr pulse, the S protein was apparent as a single gp170 component that was Endo H sensitive (lanes 1 and 2). After a 2-hr chase, a new species (gp180) was present along with gp170 in approximately equal proportions (lane3). After a 4-hr chase, the gp180 species was the dominant S protein component (lane 5) that was now Endo H resistant (lanes 5 and 6). This data is consistent with gp170 being an ER-resident glycoprotein containing high mannose chains and with gp180 corresponding with a Golgi-processed glycoprotein containing Endo H-resistant complex oligosaccharides.

The Endo H sensitivity of the C-terminus deleted $S\Delta$ protein purified from cell culture media ws also tested. As shown in Figure 76, the $S\Delta$ observed within cell lysates was found to be Endo H sensitive (lanes 1 and 2), while the secreted $S\Delta$ in cell culture media was Endo H resistant (lanes 3 and 4). This result is consistent with this glycoprotein being synthesized in an

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immature form in the ER prior to transfer to the Golgi where the complex carbohydrate is added and the protein then secreted.

As already described, the S protein expressed in COS7 cells was detected as a gp170/gp180 doublet in western blot analyses of cell lysates that were fully denatured by boiling in the presence of DTT. However, the majority of S protein was detected as a high molecular glycoprotein in the 440-669 kDa range when the same cell lysate was not heat-denatured prior to western blot analysis using SDS-PAGE (Fig. 77, lane 1). The ~500 kDa species was resistant to 10 mM DTT treatment (lane 3) and not dissociated into the monomeric form unless the lysate was first heat-denatured at 100°C (lane 4). In contrast, oligomeric form of a test protein (Thyroglobulin) of which quaternary structure is held by disulfide-linkage was converted into subunit form by the 10 mM DTT treatment. These data suggest that the ~500 kDa oligomeric form of S protein is not disulfide-linked and is heat labile. To confirm the heat-sensitivity of the ~500 kDa species of S protein, the heat-denaturation experiment was repeated but without DTT. As shown in Figure 78, heat denaturation of ~500 kDa protein at 100°C alone was sufficient to convert it into gp170/180 monomeric forms (lane 4). Using a 80°C heat-denaturation step, both the ~500 kDa and monomeric forms were detectable in similar proportion (lane 3).

In order to investigate further whether this ~500 kDa species represents an S protein oligomer in native conformation, comparative analyses with virion-derived S glycoprotein derived from Vero cell cultures was performed. The purified virious were solubilised in 1% SDS prior to Western blot analyses after SDS PAGE. The presence of the ~500 kDa spike protein oligomer was confirmed in virion particles (Fig. 79, lane 1). In addition, heat denaturation of solubilised virious produced the same oligomer-to-monomer conversion as seen with the fulllength recombinant S (lanes 2,3). The oligomeric nature of virion S was further analysed in a cross-linking experiment. Aliquots of inactivated virion from sucrose gradient fractions were treated with 10% SDS at 1% final concentration and diluted 2-fold with 0.2M Triethanolamine-HCl (pH 8, Sigma); Dimethyl suberimidate (DMS; Pierce Chemical Co.) was then added from a freshly prepared solution (10mg/ml in 0.2M Triethanolamine-HCl) at 3.3mg/ml final concentration. After 2 hr at room temperature, samples were concentrated with Centricon-30 and analyzed by silver staining after electrophoresis on a 4% polyacrylamide gel. Both untreated and DMS cross-linked virion proteins were heat-denatured, and the heat effect on the maintenance of oligomer structure was analysed by SDS-PAGE and silver staining (Figure 80). In the absence of cross-linking, heat denaturation resulted in the replacement of the ~500 kD spike protein species with the monomer species. In contrast, in the cross-linked proteins, the levels of the ~500 kD and monomer species did not change significantly after heating. These data support the fact that the ~500 kD protein is an oligomer of S monomer proteins that are bound non-covalently. After cross-linking and boiling, the ~500 kDa species migrated as a somewhat slower diffuse form

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than the untreated form. This mobility shift is probably due to a structural change resulting from boiling. In addition, a minor protein species of ~300 kD, which may represent a non-dissociated S dimer, could be seen.

To estimate more precisely the size of the recombinant ~500 kDa S species expressed in COS7 cells, a COS7 cell lysate containing the S protein oligomer was fractionated using size-exclusion column chromatography. The major portion of the ~500 kDa oligomer co-eluted with a 572 kDa marker protein. Taken together, these experiments suggest that the ~500 kDa S species seen in COS7 cell lysates is probably a homotrimer of the S protein monomer.

The oligomeric status of the $S\Delta$ spike protein was also examined after expression in COS7 cells. As shown in Figure 81, the recombinant $S\Delta$ proteins present in cell lysates were also detected in high molecular weight forms of ~500 kDa range when the lysate was not heated prior to SDS-PAGE and Western blot analysis (lane 1). However, the efficiency of oligomerization by intracellular $S\Delta$ protein appears to be much less (<10%) compared to that of full-length S protein under the same western analysis conditions. A heat-sensitivity test on this ~500 kDa protein showed that the $S\Delta$ oligomer was more heat labile than that of the full-length S oligomer, as demonstrated by the >90% conversion of all of the ~500 kDa species into monomeric $S\Delta$ forms at $S\Delta$ (Figure $S\Delta$), the majority of the secreted $S\Delta$ protein was found in monomeric form with the ~500 kDa species barely detectable (and only detectable when the protein was loaded in excess for Western analysis) (lane 1). At a temperature above $S\Delta$ 0°C, all secreted $S\Delta$ proteins were detected as monomers (lanes $S\Delta$ 3).

The ~500kDa protein is glycosylated, and the effect of deglycosylation on its antibody binding was investigated. The recombinant COS7 lysate was treated with PNGase F under non-denaturing condition (as described above) and analysed by western blot. As shown in Figure 83, deglycosylation did not affect the binding of anti-histidine Mab antibody to the treated S oligomer (lanes 2,3). However, it compromised the reactivity with the rabbit antisera raised against purified virus (lane 6). This antiserum binds to virion-derived S in western blot analyses only when DTT is omitted from the sample for SDS-PAGE indicating that it recognizes primarily a discontinuous, conformational epitope(s). This antisera has also been shown to have a high-titer of viral neutralizing antibodies. Its lack of binding to deglycosylated, recombinant S suggests that the carbohydrate actively contributes to the higher order, native structure of the S polypeptide oligomer.

The difference between the recombinant S and S Δ protein is the presence or absence of the TM-and Cys-rich domains at the C-terminus. This difference predicts that full-length S would be found associated with the membrane fraction while Sd would be in the soluble fraction upon lysis of transfected cells. Therefore, nSh- or nSh Δ TC-transfected cells were lysed under hypotonic conditions and the soluble cytosol fraction was separated from the insoluble

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membrane fraction by centrifugation (Figure 48). As shown in Figure 84, the S protein was found in the membrane fraction (DF) both as a \sim 500 kDa and 180/170 kDa species (lane 4) but was not detectable in the soluble cytosol fraction (AF) (lane 3). However, the truncated S Δ protein was found as a monomeric species (gp170) in both fractions (lanes 5,6). This indicates that the C-terminal TM and Cys-rich domains are required for the anchorage of the S protein to cell membrane.

The cellular location of the S and S Δ proteins in COS7 cells was analyzed by indirect immunofluorescence microscopy. At 48 hr post-transfection, cells were directly fixed with 2% paraformaldehyde without detergent for cell surface staining or treated with detergent followed by Cytofix/Cytoperm solution for intracellular staining. Fixed cells were then stained with rabbit anti-SARS sera (2BE) and FTC-conjugated antibody. The nSh-transfected cells showed foci of S protein indicative of Golgi-localisation (Figure 85A), while the nSh Δ TC-transfected cells displayed a uniform distribution of S Δ protein throughout the cytoplasm indicative of ER localisation (Figure 85B). While the complete S protein was also observed on the surface of transfected cells in unfixed cells (Figure 85D), the S Δ was undetectable on the cell surface (Figure 85E). These results indicate the role played by the TM-and Cys-rich domains in anchoring the S protein to the plasma membrane. Although the TM-region alone is likely responsible for membrane anchorage, the potential role played by the Cys-rich region remains to be determined.

The SARS recombinant full-length S protein is thus an N-linked glycoprotein with an estimated molecular weight of 170-180,000 kDa. Deglycosylation with PNGase F resulted in a polypeptide of the expected size for the uncleaved, encoded polypeptide (140kDa). Both transient and stable expression of the full-length SARS-CoV S gene in a variety of mammalian cells, including COS7, 293, BHK21, and Huh7 cell lines, consistently produced a S protein doublet (gp170/180) as detected in western blot analyses. Pulse-chase analyses of transfected cells demonstrated that the SARS CoV S protein was initially synthesized as an Endo H sensitive gp170 species followed by the gradual appearance of an Endo H resistant gp180 form, presumably as a result of the addition of complex carbohydrate within the Golgi apparatus.

The recombinant S protein was not secreted into the cell culture medium unless the Cterminal 60 amino acids containing the TM-region and the Cys-rich tail were deleted.

The quaternary structure of the full-length recombinant S protein was investigated using cross-linking treatment, heat-denaturation, and size fractionation analyses. The results data are consistent with the recombinant S protein existing as a homotrimer of ~500kDa. Similar analyses of virion-derived S yielded the same results. Such a trimeric structure has been reported for other enveloped RNA viruses: the hemagglutinin HA of influenza virus, the E1-E2 heterodimer of alphaviruses and the G protein of vesicular stomatitis virus. Incubations under reducing

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conditions indicate that the SARS-CoV S trimeric structure is non-covalently associated, and is very stable. S oligomers present in the cell lysate were shown to be resistant to reduction by 10 mM DTT, detergent treatment with 1% SDS, and heat denaturation at up to 60°C. Incubation at a temperature higher than >80°C resulted in the dissociation of the trimeric complex as evidenced by the decrease in trimer with the concomitant increase in the monomer bands. The temperature-induced appearance of the high-mannosylated gp170 (ER monomer form) as well as the complex-glycosylated gp180 (Golgi monomer form) suggests that trimerization can occur before the transport of the monomer spike protein to the medial Golgi apparatus. This is consistent with other reports for TGEV, influenza virus HA, and vesicular stomatitis virus G proteins. With these proteins, trimerization was reported to take place before addition of complex oligosaccharides in the Golgi apparatus.

The C-terminally truncated form of S was found in the cell lysate in both oligomeric and monomeric forms at a frequency of 10% and 90%, respectively. The truncated protein secreted into medium was found fully glysosylated and it was essentially all in monomeric form. We conclude that the C-terminal 60 amino acids of the S glycoprotein contains a membrane anchor region that affects the efficiency of trimerization. In S protein trimerization, it is possible that the C-terminal region is required to initiate the event and the triple-stranded coiled coil structures in the S2 stalk domain provide further stabilizing force as seen in HA oligomer of influenza virus.

EXAMPLE 24: CHO cells for Spike protein expression

CHO cell lines that stably express either the full-length or truncated SARS-CoV spike proteins were prepared. Several stably transfected CHO cell lines were obtained, and Figure 73 shows western blot data from a panel of representative clones.

EXAMPLE 25: Expression in E.coli

All SARS-CoV ORFs (Figure 17, Table 10) were cloned in the pET vector and expressed as C-terminal His-Tag fusion proteins in *E.coli*. The proteins smaller than 16KD were also expressed as N-terminal GST (Glutathione S-transferase) fusion proteins using pGEX vector.

Nsp1 and Nsp2, the two SARS-CoV proteins with proteolytic activity, were not expressed as full length proteins due to toxicity in *E.coli*. The respective genes were instead cloned in different portions in order to separate the catalytic residues (Cys833/ His994 for Nsp1; His41/Cys145 for Nsp2) in the resulting recombinant proteins: Nsp1A from nucleotides 2719-5214 of AY310120; Nsp1B from nucleotides 5218-7371; Nsp1C from nucleotide 7372-9984; Nsp2A from nucleotide 9985-10416; Nsp2B from nucleotide 10476-10902.

Nsp9 (SEQ ID NO: 9775) was divided into two portions: Nsp9A from nucleotide 13371-14756; Nsp9B from nucleotide 14757-16166.

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Matrix (M), ORF3 and ORF7 contain respectively three, two and one transmembrane domains. These proteins were expressed as deletion proteins excluding the first 100 amino acids (M and ORF3) or the first 18 amino acids (ORF7) that include the hydrophobic regions.

The cloned sequences are shown in Table 26.

A two-step strategy was used to amplify the cloned sequences. In the first step, amplification of DNA fragments containing more than one gene or single gene used sequenced cDNA as template. Eleven cDNA sequences were amplified: (1) a fragment, named amplC1, including genes coding for protein E, protein M, orf 7-8-9-10; (2) a fragment, named amplC2, including genes coding for orf 3-4; (3) a fragment, named amplC5, including genes coding for proteins Nsp12 and Nsp13; (4) Nsp11gene; (5) P28 and P65 genes; (6) Nsp1B and Nsp1C genes portion; (7) a fragment, named amplC9, including genes coding for proteins Nsp2 and Nsp3; (8) a fragment, named amplNsp4-7, including genes coding for proteins Nsp4, Nsp5, Nsp6, Nsp7 and for amplification of Nsp9A gene portion; (9) Nsp 9B gene portion and Nsp10 gene; (10) a fragment, named amplCO, including genes coding for proteins Orf11, Nucleocapsid (N) and Orf12; (11) Nsp1A gene portion. The primers used in this first step are given in Table 27:

In the second step, amplification of single genes was performed using DNA fragments from the first amplification step as templates. The primers are shown in Table 28.

Of the proteins where expression was seen, it was either in inclusion bodies (insoluble) or in a soluble form. Purification proceeded on appropriate material. Table 29 shows the molecular weight of the expressed fragments of SARS-CoV ORFs, whether they were cloned (+ or -), whether the cloned fragment was seen to be expressed (+ or -) and the form of protein which was chosen for purification.

Where a protein was a soluble His-tagged product, a single colony was streaked and grown overnight at 37°C on a LB/Amp (100 μ g/ml) agar plate. An isolated colony from this plate was inoculated into 20ml of LB/Amp (100 μ g/ml) liquid medium and grown overnight at 37°C with shaking. The overnight culture was diluted 1:30 into 1.0 L LB/Amp (100 μ g/ml) liquid medium and allowed to grow at the optimal temperature (30 or 37°C) until the OD550nm reached 0.6-0.8. Expression of recombinant protein was induced by addition of IPTG (final concentration 1.0 mM) and the culture incubated for a further 3 hours. Bacteria were harvested by centrifugation at 8000 x g for 15 min at 4°C. The bacterial pellet was resuspended in 10 ml of cold buffer A (300 mM NaCl, 50 mM phosphate buffer, 10 mM imidazole, pH 8.0). Cells were disrupted by sonication (or French Press) on ice four times for 30 sec at 40W using a Branson sonifier 450 and centrifuged at 13 000xg for 30 min at 4°C. Supernatants were mixed with 150 μ l Ni²⁺-resin (previously equilibrated with buffer A) and incubated at room temperature with gentle agitation for 30 min. The resin was Chelating Sepharose Fast Flow (Pharmacia), prepared according to the manufacturer's protocol. The batch-wise preparation was centrifuged at 700 x g for 5 min at 4°C

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and the supernatant discarded. The resin was washed twice (batch-wise) with 10ml buffer A for 10 min, resuspended in 1.0 ml buffer A and loaded onto a disposable column. The resin continued to be washed with buffer A at 4°C until the OD280nm of the flow-through reached 0.02-0.01. The resin was further washed with cold buffer B(300 mM NaCl, 50 mM phosphate buffer, 20 mM imidazole, pH 8.0) until the the OD280nm of the flow-through reached 0.02-0.01. The His-fusion protein was eluted by addition of 700μ l of cold elution buffer C (300 mM NaCl, 50mM phosphate buffer, 250 mM imidazole, pH 8.0) and fractions collected until the OD_{280nm} indicated all the recombinant protein was obtained. 20μ l aliquots of each elution fraction were analyzed by SDS-PAGE. Protein concentrations were estimated using the Bradford assay.

Where a protein was seen as an insoluble product, the inclusion bodies were purified as follows: homogenize cells (5 g wet weight) in 25 ml 0.1M Tris HCl pH 7, 1mM EDTA, at 4°C using an ultraturrax (10000 rpm); add 1.5mg lysozyme per gram cells; mix shortly with an ultraturrax and incubate at 4°C for 30'; use sonication or high-pressure homogenization to disrupt the cells; to digest DNA, add MgCl₂ to a final concentration of 3mM and DNase to a final concentration of 10ug/ml and incubate 30' at 25°C. add 0.5 vol of 60mM EDTA, 6% Triton x-100, 1.5M NaCl pH 7.0 to the solution, and incubate for 30' at 4°C; centrifugation at 31000 g for 10' at 4°C; re-suspend pellet in 40ml of 0.1M Tris HCl pH 7.0, 20 mM EDTA using ultraturrax; centrifugation at 31000 g for 10' a 4°C; store the IB pellet at – 20°C.

The results of expression are shown in Figures 86 to 105. Examples of purity and yield are given in Table 30.

EXAMPLE 26: Retention of critical epitope on truncated Spike antigen

A human monoclonal antibody having neutralizing activity was tested in an ELISA assay for reactivity with the purified truncated Spike protein. Briefly, ELISA plates were coated with truncated form of the spike protein at a concentration of $1\mu g/ml$ ($100\mu l/$ well) by incubating the plates overnight at 4°C. The plates were washed, non-specific binding sites were blocked and then different dilutions of the antibody were added and plates were incubated for 1 hour at room temperature. At the end of incubation, the plates were washed and bound antibody was detected by using anti-human IgG conjugated to horse radish peroxidase (HRP) and an appropriate substrate. The optical density of each well was recorded at 405 nm'using an ELISA reader. The data are shown in Figure 69 and clearly demonstrate that the neutralizing epitope recognized by the mAb is preserved and exposed on the recombinant truncated Spike protein.

EXAMPLE 27: Different Spike vaccines

Purified truncated spike protein was used to immunize mice and the level of binding antibodies induced against the truncated spike protein was determined by ELISA assay. Briefly a group of 10 mice were immunized with $3\mu g$ of truncated spike protein adjuvanted in MF59 at 0,

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4 and 8 weeks intervals. Sera samples were collected from these animals and assayed for antibodies induced by truncated spike protein in an ELISA assay. An additional group of 8 mice was immunized with 75 μ g of DNA encoding the truncated form of the spike protein on PLG particles at 0, 4 and 13 weeks intervals, the sera were collected and analyzed as above for antispike antibodies as above

The profile of binding antibodies induced in each group was plotted as geometric mean titer (GMT). Compared to a plasmid DNA vaccine expressing truncated spike antigen and delivered using a PLG microparticle formulation, the purified truncated spike protein was significantly more potent for inducing strong antibody responses. Further comparison with the antibody responses induced by inactivated BPL-SARS-CoV (already shown protective) in the same mouse strain revealed that the magnitude of antibody responses induced by purified truncated spike protein and the inactivated virus vaccine are in the same range (Figure 70).

The neutralization potential of antibodies induced by the recombinant truncated spike protein, or plasmid DNA expressing the same spike antigen, were also evaluated. The GMT values obtained in both groups are shown in Figure 71. From these data, it appears that the purified protein is significantly more effective at inducing neutralizing antibody responses against the SARS-CoV spike. Furthermore, the neutralization titers typically induced by the purified truncated spike protein are comparable to neutralization titers induced by an inactivated SARS-CoV vaccine.

Figure 72 shows a comparison of antibody binding levels (ELISA, X-axis) with neutralization titers (Y-axis). In general there is a very good correlation between the binding and neutralizing antibodies. The bottom-left grouping shows ratios 2 weeks post-3rd immunization with the DNA vaccine; the top-right grouping shows ratios 2 weeks post-2nd immunization with the protein vaccine. Both forms of vaccine show a consistent correlation.

In further experiments, the ability of a DNA vaccine to invoke an immune response in mice was studied. Mice were immunized with pCMV-nSdTC plasmid, either free or with PLG microparticles. Serum from the mice was then used as the staining antibody against cultured 293 cells that had been transfected with spike, either full-length or truncated. The cells were centrifuged prior to testing and the pellet was lysed. The antibody was tested against the culture supernatant and against the cell lysate. As shown in Figure 112, the mouse serum was able to detect spike protein in the lysate of cells that expressed full-length spike and in the supernatant of cells that expressed the truncated spike protein. Results were comparable to the staining seen when using rabbit serum that had been obtained after immunization with whole killed virus. Thus anti-spike antibodies can be induced by the use of DNA vaccination.

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EXAMPLE 28: Expression cassettes in pCMV

The sequence of plasmid pCMVKm2 is given as SEQ ID NO: 9923. Genes encoding the spike protein either in full-length form (pCMVKm2 SARS Spike nS; SEQ ID NO: 9921) or in its Δ TC form (pCMVKm2 SARS Spike nS Δ TC; SEQ ID NO: 9922) were inserted into this basic vector.

Mice were immunized with these vectors, and with similar vectors encoding the N, M or E proteins. Vectors encoding the same proteins but with optimized codon usage were also prepared. Codons were optimized for efficient human expression starting from the FRA sequence (GenBank: AY310120). The optimized sequences are: N (SEQ ID NO: 9924); M (SEQ ID NO: 9925); E (SEQ ID NO: 9926).

After administration, expression of proteins could be detected by immunofluorescence in all cases. For example, Figure 106 shows immunofluorescence (using anti-SARS rabbit serum) results after administration of the vector encoding optimsed N antigen, revealing high level expression. Mice receiving the control vector alone showed no fluorescence.

Figure 107 compares immunofluorescence (using Abgent anti-M antibody) of the native M sequence (107A) or the codon-optimsed M sequence (107B). Similarly, Figure 108 compares immunofluorescence (using Abgent anti-E antibody) of the native E sequence (108A) or the codon-optimsed E sequence (108B).

Four groups of mice (8 mice per group) were immunized with: (1) SARS nS Spike, nSdTC truncated Spike, and N proteins; (2) pCMV-SARS-nSdTC: DNA+DNA-PLG at weeks 0,4 and 13 wks; (3) CMV-nS: DNA+DNA-PLG+VEE/SIN Rep at 0, 4 and 9 wks; (4) VEE/SIN Rep-SARS-nS three times at 0, 4 and 13 wks. Sera from all groups recognized SARS nS and nSdTC proteins, and also showed virus binding and neutralization activity.

EXAMPLE 29: Spike protein cleavage

To investigate the effect of proteolytic cleavage on SARS-CoV Spike protein, it was expressed in various forms in *E.coli*, including: (1) full-length S1-S2; (2) S1 alone; (3) HR1 heptad; and (4) HR2 heptad. The expressed proteins were used to raise immune rabbit sera, which were then used for visualizing western blots of Vero cells, either infected or not infected with SARS-CoV.

Figure 109 shows a western blot using a 1:10000 dilution of antibody raised against either the S1 domain or the uncleaved S1-S2 domains. Figure 110 shows a western blot using a 1:10000 dilution of antibody raised against each of the four proteins. The difference in antigen reactivity is readily apparent.

Figure 111 shows similar data. Each serum was tested against four lanes, with those gour lanes being from left to right: (a) serum at 1:500 dilution, SARS-CoV-infected cells; (b) serum at

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1:500 dilution, non-infected cells; (c) serum at 1:2500 dilution, SARS-CoV-infected cells; (d) serum at 1:2500 dilution, non-infected cells. Again, the difference in antigen reactivity is readily apparent.

Figures 109-111 show that the Spike protein exists in various forms in infected Vero cells, with sizes of approx. 75kDa, 90kDa, 180kDa and >250kDa. The Spike protein is cleaved (at least partially) either intracellulary or after release of the particles.

If enzymatic cleavage of the mouse hepatitis coronavirus spike protein is inhibited then cell-cell fusion (syncytia formation) is also inhibited, but virus-cell fusion is not (de Haan et al. (2004) J Virol). Syncytia are observed in vivo in the lungs of SARS-infected patients, but are not seen in Vero cell cultures of the SARS-CoV. Inhibition of Spike protein cleavage may thus be used to prevent syncytia formation and related pathology, even though viral infectivity may not be blocked.

Example 30: Purification of SARS protease

Cells were grown at 37°C to mid-log phase and induced with 0.2% L-arabinose. Cells were harvested by centrifugation, and the cells resuspended in lysis buffer (LB) containing 20 mM Tris pH 7.5, 500 mM NaCl, 5% glycerol V/V, 0.05% Triton X-100, 5 mM BME, 5 mM imidazole, and complete protease inhibitors (-)EDTA. Benzonase was added to a final concentration of 50U/ml of lysate. Cells were then lysed using two passes through a pre-chilled microfluidizer. The lysate was clarified by high speed centrifugation at 44,000 x g. Clarified lysate was applied to a prepared Pharmacia chelating FF column charged with nickel sulfate. After application of the lysate the column was washed with 5 column volumes of LB. followed by 5 column volumes of LB supplemented with 45 mM imidazole. The column was then eluted using LB supplemented with 250 mM imidazole. Purity of the isolated SARS protease was 50%. Fractions containing protease were pooled, adjusted to 5 mM EDTA, and then applied to a Superdex 200 gel filtration column equilibrated in 20 mM Tris pH 7.5, 150 mM NaCl, 5% V/V glycerol, 0.05 % Triton X-100, and 5 mM DTT. Purity of the isolated SARS protease was 70%. Again, fractions containing the protease were pooled, and then stored at -80°C until used. Activity assay, mass spectrometry and western blot analysis were used to positively identify the protein (FIG 133). All steps were carried out with pre-chilled buffers, and kept at 4°C for as much of the preparation as possible.

Western of SARS Protease Purification Fractions

Protocol: Briefly, protein concentration was based on Absorbance at 280 nm, and coomassie stained gel estimates of purity. Protein was run on a 4-20% gradient gel, and transferred to nitrocellulose. The blot was then blocked with 3% BSA, probed with Mouse IgG anti-pentaHis,

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and then probed with a secondary antibody to Mouse IgG conjugated with HRP. The blot was visualized using an ECL kit (Pharmacia Biotech). The results are shown in Figure 133 where A is the sizing column pool loaded at 50, 100 and 200 ng of target protein and B is the immobilized metal affinity column pool loaded at 50, 100 and 200 ng of target protein.

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Example 31: Continuous Fluorescence Resonance Energy Transfer (FRET) Enzyme Assay The peptide containing EDANS, the fluorescence donor, and DABCYL, the fluorescence quencher (DABCYL-VNSTLQ ∇SGLRK-EDANS) was synthesized by Syn. Pep. (Dublin, CA). The peptide contains the cleavage site Gln-Ser in the middle. Meyers, G. et al. Handbook of Proteolytic Enzymes and Barrett, A et al. Academic Press, London, 1998, 726-728. The proteolytic activity of SARS protease was followed kinetically by measuring the level of formation of cleaved product that contains the fluorescence donor, SGLRK-EDANS using the Hitachi fluorometer (F-4500 FL Spec.) set at 340 nm excitation and 490 nm emission wave length. 5 μL of 5 mM peptide stock in DMSO solution was added to the reaction mixture, containing 295 μl of standard buffer (75 mM Tris-Hcl, 25 mM NaOAc, 25 mM Bis-Tris, 25 mM glycine, 5 mM EDTA, and 1 mM EDTA, pH 7.4) and 100 ul of buffer or 100 ul of 3.6 uM protease stock solution. The kinetic curve was followed for 6 minutes (the reaction was linear with R2 value of 0.998 (FIG 134)). The formation of fluorescence (proteolytic reaction) is likely enzyme dependent, as concentration of enzyme was tripled three times as much fluorescence was formed in the 6 minutes time frame.

It will be understood that the invention has been described by way of example only and modifications may be made whilst remaining within the scope and spirit of the invention.

Table 1. US Patents and Published International Patent Applications

Publication		Publication
Number		Date
		12/16/1975
		3/1/1977
		12/27/1977
		5/16/1978
US-4122191		10/24/1978
		3/11/1980 -
		3/3/1981
US-4264617	Antiviral 5-(Substituted Benzal) Hydantoins	4/28/1981
		9/1/1981
US-4327088	Phosphonooxy- Or Glycosyloxy-Substituted Acrylophenones, Compositions And Uses Thereof	4/27/1982
US-4332820	Substituted Benzonitriles Having Antiviral Activity	6/1/1982
US-4349568	Sulfur-Substituted Diphenyl Ethers Having Antiviral Activity	9/14/1982
US-4352792	3-Alkoxyflavone Antiviral Agents	10/5/1982
US-4371537	Sulfur-Substituted Phenoxypyridines Having Antiviral Activity	2/1/1983
		12/27/1983
	Process For The Preparation Thereof	
US-4505929	Sulfur-Substituted Diphenyl Ethers Having Antiviral Activity	3/19/1985:
US-4526897		7/2/1985
US-4558134	Certain Phenoxy-Pyridine-Carbonitriles Having Antiviral Activity	12/10/1985
	Endowed With Anti-Viral Activity 2-Alkylamino-4,6-Dihalo Pyrimidines	12/16/1986
US-4636492	Inhibition Of Viral Protease Activity By Peptide Halomethyl Ketones	1/13/1987
US-4652552	Tetrapeptide Methyl Ketone Inhibitors Of Viral Proteases	3/24/1987
US-4724233		2/9/1988
		4/19/1988
US-4847246		7/11/1989
US-4855283		8/8/1989
	And -Ureas	
	Phosphorus Compounds, Processes For Their Manufacture, And Their Use	12/5/1989
US-4956351	Antiviral Pharmaceutical Compositions Containing Cyclodextrins	9/11/1990
		3/19/1991
US-5036072	Antiviral Agent	7/30/1991
US-5070090	Antipicorpaviral Herterocyclic-Substituted Morpholinyl Alkylphenol Ethers	12/3/1991
US-5100893	Antipicomaviral Pyridazinamines	3/31/1992
US-5112825	Antirhinoviral Heteroamine-Substituted Pyridazines	5/12/1992
US-5157035	Anti-Virally Active Pyridazinamines	10/20/1992
US-5240694	Combined Antiviral And Antimediator Treatment Of Common Colds	8/31/1993
US-5242924	Tetrazolyl-(Phenoxy And Phenoxyalkyl)-Piperidinylpyridazines As Antiviral Agents	9/7/1993
US-5278184	Synthetic Derivatives Of Pyrrole And Pyrrolidine Suitable For The Therapy Of Infections Caused	1/11/1994
	By Rhinoviruses	
US-5364865	Phenoxy- And Phenoxyalkyl-Piperidines As Antiviral Agents	11/15/1994
US-5453433	Thiadiazoles And Antipicornaviral Compositions	9/26/1995
US-5492689	Combined Virustatic Antimediator (COVAM) Treatment Of Common Colds	2/20/1996
US-5514679	Therapeutic Phenoxyalklpyridazines And Intermediates Therefor	5/7/1996
US-5514692	Substituted Quinoline Derivatives Useful As Antipiconaviral Agents	5/7/1996
US-5523312	Antipicornaviral Agents	6/4/1996
US-5545653	Anti-Viral Compounds	8/13/1996
US-5552420	Therapeutic Phenoxyalkylazoles And Phenoxyalkylazines	9/3/1996
	Thiadiazoles And Their Use As Antipicornaviral Agents	10/22/1996

PCT/US2004/011710

US-5580897	1,2-Dithiins Having Antifungal Activity	12/3/1996
US-5618821	Therapeutic Phenoxyalkylheterocycles	4/8/1997
US-5618849	Orally Active Antiviral Compounds	4/8/1997
US-5648354	1,2-Dithiins Having Antifungal Activity	7/15/1997
	Thiadiazoles And Their Use As Antipicornaviral Agents	7/22/1997
US-5693661	Anti-Viral Compounds	12/2/1997
US-5721261	Therapeutic Phenoxyalkylazoles And Phenoxyalkylazines	2/24/1998
US-5725859	Plant-Based Therapeutic Agent With Virustatic And Antiviral Effect	3/10/1998
US-5750527	Thiadiazoles And Their Use As Antipicornaviral Agents	5/12/1998
US-5750551	Treatment For Viral Diseases	5/12/1998
US-5762940	Methods And Compositions For Inhibiting Or Destroying Viruses Or Retroviruses	6/9/1998
US-5763461	Therapeutic Phenoxyalkylheterocycles	6/9/1998
US-5821242	Anti-Viral Compounds	10/13/1998
US-5821257	Thiadiazoles And Their Uses As Antipicomaviral Agents	10/13/1998
US-5821331	Anti-Picornaviral Agents	10/13/1998
	Therapeutic Phenoxyalkylazoles And Phenoxyalkylazines	12/8/1998
US-5856530	Antipicomaviral Compounds And Methods For Their Use And Preparation	1/5/1999
US-5891874	Anti-Viral Compound	4/6/1999
US-5962487	Antipicornaviral Compounds And Methods For Their Use And Preparation	10/5/1999
US-6004933	Cysteine Protease Inhibitors	12/21/1999
US-6020371	Antipicomaviral Compounds Compositions Containing Them And Methods For Their Use	2/1/2000
US-6087374	Anti-Viral Compounds	7/11/2000
US-6114327	Anti-Viral Compounds	9/5/2000
US-6117844	Method And Composition For Antiviral Therapy	9/12/2000
US-6194447	Bis (Benzimidazole) Derivatives Serving As Potassium Blocking Agents	2/27/2001
US-6214799		4/10/2001
US-6277891	Nitric Oxide Inhibits Rhinovirus Infection	8/21/2001
US-6294186	Antimicrobial Compositions Comprising A Benzoic Acid Analog And A Metal Salt	9/25/2001
US-6331554	Antipicomaviral Compounds, Compositions Containing Them, And Methods For Their Use	12/18/2001
US-6358971	Anti-Viral Compounds	3/19/2002
US-6362166	Antipicornaviral Compounds And Methods For Their Use And Preparation	3/26/2002
US-6414004	3-Substituted 5-Aryl-4-Isoxazolecarbonitriles Having Antiviral Activity	7/2/2002
US-6420591	Carbamates And Compositions Thereof, And Methods For Their Use For Treating Cancer,	7/16/2002
	Inflammation, Or A Viral Infection	1012002
US-6469018	Compounds	10/22/2002
US-6498178	Inhibitors Of IMPDH Enzyme	12/24/2002
US-6514997	Antipicornaviral Compounds And Compositions, Their Pharmaceutical Uses, And Materials For	2/4/2003
ı	Il neir Synthesis	1,2003
US-6525043	Use Of Ion Channel Modulating Agents	2/25/2003
US-6531452	Antipicornaviral Compounds And Compositions, Their Pharmaceutical Uses, And Materials For	3/11/2003
	Their Synthesis	
US-6534489	Organophosphorus Compounds And The Use Thereof	3/18/2003
WO00/06529	Diketoacid-Derivatives As Inhibitors Of Polymerases	2/10/2000
WO00/25791	Pyridin-4-Yl Or Pyrimidin-4-Yl Substituted Pyrazines	5/11/2000
WO00/27423	Methods And Compositions For Treating Common Cold Symptoms	5/18/2000
WO00/34308	Protein Transduction System And Methods Of Use Thereof	6/15/2000
WO00/39348	Methods And Compositions For Identifying Protease Modulators	7/6/2000
WO00/40243	Novel Compounds	7/13/2000
WO00/50037	Nitrosated And Nitrosylated Proton Pump Inhibitors, Compositions And Methods Of Use	8/31/2000
IWO00/56331	Inhibitors Of Impdh Enzyme	0.000.000
WO00/56757	Immunomodulatory Steroids. In Particular The Hemibydrate Of 16 Alpha Bromogniandsostoropa	9/28/2000
W 0000/66096	New Indication For Use Of Antiepileptic Agents And Medicines	11/9/2000
WO00/78746		12/28/2000

WOO1/0535 Pyrazolidinol Compounds 1/4/2001 1/1/20		Compounds Obtained From Salvia Species Having Antiviral Activity	1/4/2001
WOO1/03581 Urus Like Particles, Their Preparation And Their Use Preferably In Pharmaceutical Screening And Functional Genomics WOO1/03681 Use Of Flavones, Coumarins And Related Compounds To Treat Infections WOO1/03681 Use Of Cobalt Chelates For Treating Or Preventing Virus Infection WOO1/03681 Antipicomaviral Compounds And Compositions, Their Pharmaceutical Uses, And Materials For Their Synthesis WOO1/19322 Use Of Cosals In Rhinovirus Infection WOO1/19322 Ise of Cosalsd In Rhinovirus Infection WOO1/19322 Ise of Cosalsd In Rhinovirus Infection WOO1/19322 Antiviral Agents WOO1/272920 Colon And Colon Cancer Associated Polynucleotides And Polypeptides WOO1/273185 Novel Carbamates And Ureas WOO1/31016 Processed Human Chemokines Pho-1 And Pho-2 WOO1/37312 3,4-Dihydro-(Ih)-Quinazolin-2-Ones And Their Use As Csbp/P38 Kinase Inhibitors WOO1/378373 3,4-Dihydro-(Ih)-Quinazolin-2-One Compounds As Csbp/P38 Kinase Inhibitors WOO1/38313 3,4-Dihydro-(Ih)-Quinazolin-2-One Compounds As Csbp/P38 Kinase Inhibitors S731/2001 WOO1/38313 3,4-Dihydro-(Ih)-Quinazolin-2-One Compounds As Csbp/P38 Kinase Inhibitors S731/2001 WOO1/38313 3,4-Dihydro-(Ih)-Quinazolin-2-One Compounds As Csbp/P38 Kinase Inhibitors S731/2001 WOO1/38013 3,4-Dihydro-(Ih)-Quinazolin-2-One Compounds As Csbp/P38 Kinase Inhibitors S731/2001 WOO1/38013 3,4-Dihydro-(Ih)-Quinazolin-2-One Compounds As Csbp/P38 Kinase Inhibitors S731/2001 WOO1/40189 Antipicomaviral Compounds And Compositions, Their Pharmaceutical Uses, And Materials For Their Synthesis WOO1/40303 Multivalent Electron Active Compositions And Methods Of Making And Using Same 7/12/2001 WOO1/79167 Antipicomaviral Compounds And Compositions, Their Pharmaceutical Uses, And Materials For Their Synthesis WOO1/90129 Prophylactic And Therapeutic Treatment Of Infectious And Other Diseases With Mono-And Diseascharide Based Compounds WOO1/93883 Therapeutic Agents - Ii 1/27/2001 WOO1/938	WO01/00585	Pyrazolidinol Compounds	
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WO03/20222	Dioxolane And Oxathiolane Derivatives As Inhibitors Of Pna Dependent Pna Visal Palameter	3/13/2003
WO03/20270		3/13/2003
	Anti-Picornaviral Agents	5/13/2003
WO03/20271	Oxadiazolyl-Phenoxyalkylisoxazoles, Compositions Thereof And Methods For Their Use As	3/13/2003
	Anti-Picornaviral Agents	5/13/2003
WO03/20712	Oxadiazolyl-Phenoxyalkylthiadiazoles, Compositions Thereof And Methods For Their Use As	3/13/2003
1	Anti-Picornaviral Agents	D/13/2003
WO86/03412	Improvements Relating To The Treatment Control And Prevention Of Rhinovirus Infections	6/19/1986
WO86/039/1	Antiviral Agents	7/17/1986
WO88/09669	Avirulent Microbes And Uses Therefor	12/15/1988
WO92/03475	Enterovirus Peptides	3/5/1992
WO92/22520	Orally Active Antiviral Compounds	12/23/1992
WO92/22570	Inhibitors Of Picornavirus Proteases	12/23/1992
WO94/00012	Nucleic Acids And Methods Of Use Thereof For Controlling Viral Pathogens	1/6/1994
WO95/03821	Prosaposin And Cytokine-Derived Peptides As Theraneutic Agents	2/9/1995
WO95/09175	Ring-Expanded Nucleosides And Nucleotides	4/6/1995 ·-
WO95/11992	Antiviral Compounds	5/4/1995
WO95/31198	Thiadiazoles And Their Use As Antinicornaviral Agents	11/23/1995
WO95/31438	Therapeutic Phenoxyalkylheterocycles	11/23/1995
WO95/31439	Therapeutic Phenoxyalkylnyridazines And Intermediates Therefor	11/23/1995
WO95/31452		11/23/1995
WO95/34595	Antiviral Dendrimers	12/21/1995
WO95/35103		12/28/1995
1	Optionally Inflammations As Well As A Method For The Treatment Thereof	12/20/1993
WO96/05836	Methods Of Treating Cold Symptoms Heing Pentovifulling	2/29/1996
WO96/05854		2/29/1996
	Denvative	42311330
WO96/09822	Antipicornaviral Agents	1/4/1996
WO96/11211	Selective Inhibition Of Internally Initiated Rna Translation	1/18/1996
WO96/22689	Multiple Component Rna Catalysts And Uses Thereof	3/1/1996
WO96/40641	Sulfonamide Derivatives As Cell Adhesion Modulators	12/19/1996
WO97/08553	Targeting Of Proteins To The Cell Wall Of Gram-Positive Bacteria	
WO97/34566	Electrophilic Ketones For The Treatment Of Hernesvirus Infections	3/6/1997
WO97/41137	Use Of Anthocyanidin And Anthocyanidin Derivatives	0/25/1997 1/6/1997
WO97/43305	Inhibitory Of Discourse 2 D	1/20/1997
	I among the first of the repairment	1120/1991

	Novel Anti-Viral Compounds	12/18/1997
	Antiviral Linear Polymers	1/29/1998
WO98/07745	Compositions And Methods For Treating Infections Using Analogues Of Indolicidin	2/26/1998
WO98/11778	Antimicrobial Treatment For Herpes Simplex Virus And Other Infectious Diseases	3/26/1998
	Antikinin Compounds And Uses Thereof	5/28/1998
	Anti-Viral Compounds	7/23/1998
WO98/31374	Method Of Treating Rhinoviral Infections	7/23/1998
	Therapeutic Treatment And Prevention Of Infections With A Bioactive Material Encapsulated Within A Biodegradable-Biocompatible Polymeric Matrix	7/30/1998
WO98/34601	Method For Inhibiting Intracellular Viral Replication	8/13/1998
WO98/42188	Antimicrobial Prevention And Treatment Of Human Immunedeficiency Virus And Other	10/1/1998
	Infectious Diseases	
WO98/43950	Antipicornaviral Compouds, Compositions Containing Them, And Methods For Their Use	10/8/1998
WO98/49190	Substituted Oxadiazole Cysteine Protease Inhibitors	11/5/1998
	Anti-Viral Compounds	12/10/1998
WO99/30699	Modulators Of Cysteine Protease	6/24/1999
WO99/31122	Antipicornaviral Compounds And Methods For Their Use And Preparation	6/24/1999
	Cysteine Protease Inhibitors	10/28/1999
	Inhibitors Of Impdh Enzyme	11/4/1999
WO99/57135	Antipicornaviral Compounds, Their Preparation And Use	11/11/1999
	Anti-Viral Compounds	11/25/1999
WO99/61437	Novel 2-Alkyl Substituted Imidazole Compounds	12/2/1999

Table 2. US Patents and Published International Patent Applications

Publication Number	mber						
WO02/69903	Nucleosides, Preparation Thereof And Use As Inhibitors Of Rna Viral Polymerases	n Date 9/12/2002					
WO02/48116	Inhibitors Of Hepatitis C Virus Ns3 Protease	6/20/2002					
WO02/48157	Imidazolidinones And Their Related Derivatives As Hepatitis C Virus Ns3 Protease Inhibitors	6/20/2002					
WO02/61048	In Vitro System For Replication Of Rna-Dependent Rna Polymerase (Rdrp) Viruses	8/8/2002					
WO03/02518	Novel 2,4-Difluorobenzamide Derivatives As Antiviral Agents	1/9/2003					
WO02/79187	Methoxy-1,3,5-Triazine Derivatives As Antiviral Agents	10/10/2002					
WO01/78648	6-Methylnicotinamide Derivatives As Antiviral Agents	10/25/2001					
WO01/12214	MYCOPHENOLATE MOFETIL IN ASSOCIATION WITH PEG-IFN-Alpha.	2/22/2001					
	4'-Substituted Nucleosides	12/19/2002					
WO02/18404	Nucleoside Derivatives	3/7/2002					
WO02/94289	Antiviral Nucleoside Derivatives	11/28/2002					
WO96/39500	Oligonucleotides Specific For Hepatitis C Virus	12/12/1996					
WO03/00713	Nucleoside Compounds In Hcv	1/3/2003					
WO01/60381	Nucleoside Analogs With Carboxamidine-Modified Bicyclic Base	8/23/2001					
WO02/03997	Pyrido[2,3-D]Pyrimidine And Pyrimido[4,5-D]Pyrimidine Nucleosides	1/17/2002					
WO97/26883	Modulation Of Th1/Th2 Cytokine Expression By Ribavirin3 And Ribavirin3 Analogs In Activated T-Lymphocytes	7/31/1997					
WO03/26589	Methods And Compositions For Treating Hepatitis C Virus Using 4'-Modified Nucleosides	4/3/2003					
WO03/26675	Methods And Compositions For Treating Flaviviruses And Pestiviruses Using 4'-Modified Nucleoside	4/3/2003					
WO97/30067	Sugar-Modified Gapped Oligonucleotides	8/21/1997					
WO01/47883	Fused-Ring Compounds And Use Thereof As Drugs	7/5/2001					
	Fused Cyclic Compounds And Medicinal Use Thereof	1/3/2003					
	Pyrrolo[2,3-D]Pyrimidine Nucleoside Analogs	12/19/2002					
WO01/55111	Biaryl Compounds, Their Preparation And Their Use In Therapy	8/2/2001					

WO01/16149	2-Azapurine Compounds And Their Use	3/8/2001
WO01/85770	Sentinel Virus Ii	11/1/10001
WO02/12263	Nucleic Acid Binding Compounds Containing Pyrazolo[3,4-D]Pyrimidine Analogues Of Purin- 2,6-Diamine And Their Uses	2/14/2002
JP 2001-247550 A	2 Condensed Ring Compound And Its Medicinal Use	
6210675	PT-NANB Hepatitis Polypeptides	9/11/2001
6451991	Sugar-Modified Gapped Oligonucleotides	4/3/2001
5830455	Method Of Treatment II-in A 77	9/17/2002
	Method Of Treatment Using A Therapeutic Combination Of α-Interferon And Free Radical Scavengers	11/3/1998
5908621	Polyethylene Glycol Modified Interferon Therapy	6/1/1999
5990276	Synthetic Inhibitors Of Hepatitis C Virus NS3 Protease	11/23/1999
6172046	Combination Therapy For Eradicating Detectable HCV-RNA In Patients Having Chronic Hepatitis C Infection	1/9/2001
6177074	Polyethylene Glycol Modified Interferon Therapy	1/23/2001
6326137	Hepatitis C Virus Protease-Dependent Chimeric Pestivings	
6434489	Compositions Of Hepatitis C Virus NSSR Polymerase And Methods For Countablishing St.	12/4/2001
6461605	Continuous Low-Dose Cytokine Infusion Therapy	8/13/2002
6472373	Combination Therapy For Fradition Post and average and a second	10/8/2002
	Patients Having Chronic Hepatitis C Infection	10/29/2002
6524570	Delustrates Class 136 VC 17 C	
WO00/37097	Ribavirin Interferon Alfa Industria IV. C. 1: .: PR	2/25/2003
WO00/37110	Ribavirin Pegulated Interferen Alfa I. J. J. Y. C. 11	6/29/2000
WO00/62799		6/29/2000
WO01/58929		10/26/2000
WO02/32414	Pihavirin Pagulated Interferen Alf- IV C 11 11 77	8/16/2001
WO03/24461		4/25/2002
WO93/20835	Treatment Of Handida Will C. C.S.	3/27/2003
WO96/36702	Soluble Active Wastick CVIII B	10/28/1993
WO97/16204	Continuous I on Deer Carti Y C : m	11/21/1996
WO97/43310	Synthetic Inhibitore Of II	5/9/1997
WO98/48840	Synthetic Inhibitors Of Hepatitis C Virus Ns3 Protease	11/20/1997.
WO99/15194	Polyethylene Glycol-Interferon Alpha Conjugates For Therapy Of Infection	11/5/1998
	preparitis C infection	4/1/1999
WO99/59621	INAIVE Patients Having G Chronic Henatitis C Infection	11/25/1999
WO02/100846	Compounds And Methods For The Treatment Or Prevention Of Flavining Infections	12/19/2002
WO02/100851	Compounds And Methods For The Treatment Or Prevention Of Flavining Infantions	12/19/2002
5241053	Fused Proteins Comprising Glycoprotein Gd Of HSV-1 And LTB	3/31/1993
5556946	Interleukin-2/Viral Antigen Protein Chimers	9/17/1996
6087484	Ennancement Of Ribozyme Catalytic Activity By A 2'-O-Substituted Facilitator Oligonucleotide	7/11/2000
5830905	Compounds, Compositions And Methods For Treatment Of Hepatitis C	1/3/1998
6316492		
6440985	Wethods For Treating Viral Infections	1/13/2001
WO00/10573	Compounds Compositions A-136 II 1 7 7	3/27/2002 3/2/2000
WO00/13708	Methode For Trooting O. Daniel, XV. 1 x 5	
	Methods For Treating Or Preventing Viral Infections And Associated Diseases 3	/16/2000
WO99/51781	Methods For Treating Or Preventing Viral Infections And Associated Diseases 4 Henrific C.V.: N. S. C.	/6/2000
W 099/31/61	riepatins C Virus NS5b Compositions And Methods Of Use Thereof	0/14/1999
	riepatitis C Inhibitor Tri-Peptides	1/27/2001
	riepatitis C Inhibitor Peptide Analogues	1/7/2000
	riepanus C Innibitor Tri-Peptides	2/11/2001
034941/		2/11/2001

6410531	Hepatitis C Inhibitor Tri-Peptides	6/25/2002
6420380	Hepatitis C Inhibitor Tri-Peptides	7/16/2002
6448281	Viral Polymerase Inhibitors	9/10/2002
6479508	Viral Polymerase Inhibitors	11/12/2002
6534523	Hepatitis C Inhibitor Tri-Peptides	3/18/2003
WO00/09543	Hepatitis C Inhibitor Tri-Peptides	2/24/2000
WO00/09558	Hepatitis C Inhibitor Peptides	2/24/2000
WO00/59929	Macrocyclic Peptides Active Against The Hepatitis C Virus	10/12/2000
WO02/04425	Viral Polymerase Inhibitors	1/17/2002
WO02/70739	Hcv Polymerase Inhibitor Assay	9/12/2002
WO03/07945	Viral Polymerase Inhibitors	1/30/2003
WO03/10140	Viral Polymerase Inhibitors	2/6/2003
WO03/10141	Viral Polymerase Inhibitors	2/6/2003
WO99/07734	Hepatitis C Inhibitor Peptide Analogues	2/18/1999
WO01/16379		3/8/2001
WO02/07761	Inhibiting Hepatitis C Virus Processing And Replication	1/31/2002
WO02/57287	Nucleoside Derivatives As Inhibitors Of Rna-Dependent Rna Viral Polymerase	7/25/2002
WO02/57425	Nucleoside Derivatives As Inhibitors Of Rna-Dependent Rna Viral Polymerase	7/25/2002
WO02/70651		9/12/2002;
WO03/20222	Dioxolane And Oxathiolane Derivatives As Inhibitors Of Rna-Dependent Rna Viral Polymerase	3/13/2003
PCT/US2003/	Thiosemicarbazones as Anti-Virals and Immunopotentiators	01/10/2003
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<u>Table 3</u>: US Patents and published international patent applications relating to inhalation technology for the delivery of antiviral compounds of the invention.

Publication	Title				
Number		Date			
5740794		4/21/1998			
5775320		7/7/1998			
5785049		7/28/1998;			
5814607	Pulmonary delivery of active fragments of parathyroid hormone	9/29/1998			
5826633		10/27/1998			
5458135	Method and device for delivering aerosolized medicaments	10/17/1995			
5607915	Pulmonary delivery of active fragments of parathyroid hormone	3/4/1997			
5654007	Methods and system for processing dispersible fine powders	8/5/1997			
5922354	Methods and system for processing dispersible fine powders	7/13/1999			
5928469	Process for storage of materials	7/27/1999			
5976574	Processes for spray drying hydrophobic drugs in organic solvent suspensions	11/2/1999			
5985248	Processes for spray drying solutions of hydrophobic drugs and compositions thereof	11/16/1999			
5994314	Compositions and methods for nucleic acid delivery to the lung	11/30/1999			
5997848	Methods and compositions for pulmonary delivery of insulin	12/7/1999			
6001336	Processes for spray drying aqueous suspensions of hydrophobic drugs and compositions thereof	12/14/1999			
6019968	Dispersible antibody compositions and methods for their preparation and use	2/1/2000			
6051256	Dispersible macromolecule compositions and methods for their preparation and use	4/18/2000			
6071428	Stable compositions	6/6/2000			
6077543	Systems and processes for spray drying hydrophobic drugs with hydrophilic excipients	6/20/2000			
6080721	Pulmonary delivery of active fragments of parathyroid hormone	6/27/2000			
6089228	Apparatus and methods for dispersing dry powder medicaments	7/18/2000			
6103270	Methods and system for processing dispersible fine powders	8/15/2000			
6123936	Methods and compositions for the dry powder formulation of interferons	9/26/2000			

(10/01/		
6136346	Powdered pharmaceutical formulations having improved dispersibility	10/24/2000
6138668	Method and device for delivering aerosolized medicaments	10/31/2000
6165463	Dispersible antibody compositions and methods for their preparation and use	12/26/2000
6182712	Power filling apparatus and methods for their use	2/6/2001
6187344	Powdered pharmaceutical formulations having improved dispersibility	2/13/2001
6207135	Gaseous microparticles for ultrasonic diagnosis and process for their production	3/27/2001
6231851	Methods and compositions for the dry powder formulation of interferons	5/15/2001
6257233	Dry powder dispersing apparatus and methods for their use	7/10/2001
6258341	Stable glassy state powder formulations	7/10/2001
6267155	Powder filling systems, apparatus and methods	7/31/2001
6294204	Method of producing morphologically uniform microcapsules and microcapsules produced by this method	9/25/2001
6303582	Compositions and methods for nucleic acid delivery to the lung	
6309623	Stabilized preparations for use in metered dose inhalers	10/16/2001
6309671	Stable glassy state powder formulations	10/30/2001
6358530	Powdered pharmaceutical formulations having improved dispersibility	10/30/2001
6365190	Systems and processes for spray drying hydrophobic drugs with hydrophilic excipients	3/19/2002
6372258	Methods of spray-drying a drug and a hydrophobic amino acid	4/2/2002
6423344	Dispersible macromolecule compositions and methods for their preparation and use	4/16/2002
6426210	Storage of materials	7/23/2002
6433040	Cabilized bit at	7/30/2002
6440337	Mathed and appearations and methods of use	8/13/2002
RE37872	Method and apparatus for the formation of particles Storage of materials	8/27/2002
6479049	Mothodo and commercial for the latest and a second	10/8/2002
6503411	Methods and compositions for the dry powder formulation of interferons Stable compositions	11/12/2002
6509006	Stable compositions	1/7/2003
6514496	Devices compositions and methods for the pulmonary delivery of aerosolized medicaments	1/21/2003
6518239	Dispersible antibody compositions and methods for their preparation and use	2/4/2003
6543448	dry powder compositions having improved dispersivity	2/11/2003
6546929	apparatus and methods for dispersing dry powder medicaments	4/8/2003
WO 00/15262	dry powder dispersing apparatus and methods for their use	4/15/2003
	dry powder active agent pulmonary delivery	3/23/2000
WO 93/00951	method and device for delivering aerosolized medicaments	1/21/1993
WO 94/07514	pulmonary delivery of active fragments of parathyroid hormone	4/14/1994
WO 95/24183	methods and compositions for pulmonary delivery of insulin	9/14/1995
WO 95/31479	methods and compositions for the dry powder formulation of interferons	11/23/1995
WO 96/09085	apparatus and methods for dispersing dry powder medicaments	3/28/1996
WO 96/32096	powdered pharmaceutical formulations having improved dispersibility	10/17/1996
WO 96/32116	compositions and methods for nucleic acid delivery to the lung	10/17/1996
WO 96/32149	pulmonary delivery of aerosolized medicaments	10/17/1996
WO 96/32152	pulmonary administration of dry powder alpha 1-antitrypsin	10/17/1996
WO 96/40068	methods and system for processing dispersible fine powders	12/19/1996
WO 97/41031	powder filling systems, apparatus and methods	11/6/1997
WO 97/41833	dispersible macromolecule compositions and methods for their preparation and use	11/13/1997
WO 98/16205	stable glassy state powder formulations	4/23/1998
WO 98/29096	aerosolized hydrophobic drug	7/9/1998
WO 98/29098	processes for spray drying aqueous suspensions of hydrophobic drugs with hydrophilic	7/9/1998
	excipients and compositions prepared by such processes	112/1220
WO 98/29140		7/9/1998
WO 98/29141	processes for spray drying solutions of hydrophobic drugs with hydrophilic excipients and	
	compositions prepared by such processes	7/9/1998
WO 99/19215	nowder filling amounts and and 1	
	iguid crustal forms of and	4/22/1999
	227	8/26/1999

	aerosolized active agent delivery	9/23/1999
WO 99/62495	dry powder dispersing apparatus and methods for their use	12/9/1999
WO 00/21594	flow resistance modulated aerosolized active agent delivery	4/20/2000
WO 00/61178	pulmonary administration of dry powder formulations for treating infertility	10/19/2000
WO 00/72904	apparatus and method for dispensing metered amount of aerosolized medication	12/7/2000
WO 01/00263	systems and methods for aerosolizing pharmaceutical formulations	1/4/2001
WO 01/00312	spray drying process for preparing dry powders	1/4/2001
WO 01/32144	dry powder compositions having improved dispersivity	5/10/2001
WO 01/43529	receptacles to facilitate the extraction of powders	6/21/2001
WO 01/43530	systems and methods for extracting powders from receptacles	6/21/2001
WO 01/43802	systems and methods for treating packaged powders	6/21/2001
WO 01/44764	systems and methods for non-destructive mass sensing	6/21/2001
WO 01/87393	systems, devices and methods for opening receptacles having a powder to be fluidized	11/22/2001
WO 01/93932	lockout mechanism for aerosol drug delivery devices	12/13/2001
WO 02/09669	apparatus and process to produce particles having a narrow size distribution and particles made thereby	2/7/2002
WO-02/11695	inhaleable spray dried 4-helix bundle protein powders having minimized aggregation	2/14/2002
WO 02/49619	induced phase transition method for the production of microparticles containing hydrophilic active agents	6/27/2002
WO 02/49620	induced phase transition method for the production of microparticles containing hydrophobic active agents	6/27/2002
WO 02/54868		7/18/2002
		11/7/2002
WO 02/100548		12/19/2002
	powder aerosolization apparatus and method	1/3/2003
	flow regulator for aerosol drug delivery device and methods	1/3/2003

TABLE 4: Forward and reverse primers for nucleic acid amplification of SARSV

Pair Number	Forward primer SEQ ID NO	Forward Primer Start	Forward Primer Stop	Forward Primer Tm	Forward Primer %GC	Reverse primer SEQ ID NO	Reverse Primer Start	Reverse Primer Stop	Reverse Primer Tm	Reverse Primer %GC	Primer Tm Diff	Product Length	Product Tm	Product %GC	Anneal Score	Optimum Anneal Temp
	1021	12726	12746	51.3	47.6	3521	12996	12977	50.2	40	1	271	75	42.8	26	52.6
- 2		12236	12256	51.2		3522	12993	12975	51.4	47.4	0.2	758	76.4	42.5	26	54
		12373		50.8	47.4	3523	12993	12975	51.4	47.4	0.6	621	76.4	43	26	53.8
-		12236		51.2	42.9	3524	12996	12977	50.2	40	0.9	761	76.4	42.3	26	53.6
		12373	12391	50.8	47.4	3525	12996	12977	50.2	40	0.5	624	76.4	42.8	26	53.6
	1026	12726	12746	51.3	47.6	3526	12993	12975	51.4	47.4	0.1	268	75.1	43.3	26	53.1
	10141	2671	2692	52.1	40.9	3527	3185	3164	51	45.5	1.2	515	75.6	41.6	26	53.3
	1028	28942	28961	50.2	45	3528	29298	29280	51.4	52.6	1.2	357	76.4	44.8	26	53.6
		19801	19819	53.2	52.6	3529	19922	19901	51.5	45.5	1.7	122	72.2	43.4	26	51.1
10	1030	19800		50.4	50	3530	19921	19901	50.2	47.6	0.3	122	72.2	43.4	26	50.7
11		9930		51.5		3531	10605	10588	51.1	50	0.4	676	75.8	41.3	27	53.5
12		9933	9952	50.9	45	3532	10605	10588	51.1	50	0.2	673	75.8	41.2	27	53.4
13	1033	9930	9949	52.2	50	3533	10605	10588	51.1	50	1.1	676	75.8	41.3	27	53.5

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14	1034	9927	9945	50.8	52.6	3534	10605	10588	51.1	50	0.3	679	75.8	1 44 6	T 0	T =0.4
15	1035	3789	3806	50			4445	4425	50.6		0.5	657	75.5		_	
16	1036	3788		50			4444	4424			0.5					
17		3795	3813	52.1	52.6	1-000	4445	4425	50.6		1.5		75.5			
18	1038	3787	3804	50		3538	4445	4425	50.6			651	75.5			_
19		19801	19819	53.2			19921	19900	51.8		0.5	659	75.4	40.4		
20		24418	24436								1.4	121	72.3	43.8	_	
21		9929	9949	53.8		3541	25182	25164	51.4		1.4	765	76.1	41.7		
22		2671	2692	52.1			10449	10425	54.6		0.8	521	75.4	40.9	28	54
23		3792	3810	52.1	40.9		3186	3165	50.4		1.7	516	75.6	41.5	28	53.1
24	1.4.4	9933	9952		52.6		4446	4425	51.8		1.1	655	75.5	40.6	-28	53.5
25		3792		50.9	45		10449	10431	50.9	47.4	0.1	517	75.3	40.8	28	53.1
26	1		3810	52.9		3545	4445	4424	51.3		1.6	654	75.5	40.5	28	53.3
27	1046	25782 9927	25806	53.5	40		26184	26164	52.4	42.9	1.1	403	74.7	40.2	28	53.1
28		9927	9945	50.8	52.6		10449	10431	50.9	47.4	0.1	523	75.4	40.9	28	53.1
29			9945	50.8	52.6		10449	10428	51.9	40.9	1.1	523	75.4	40.9	28	53.1
30		3789	3806	50	50		4444	4424	50.6	42.9	0.5	656	75.5	40.5	28	53
	1050	3795	3813	52.1	52.6		4444	4424	50.6	42.9	1.5	650	75.5	40.6	28	53.1
	1051	9933	9952	50.9	45		10449	10428	51.9	40.9	1.1	517	75.3	40.8	28	53.1
	1052	9930	9948	51.5	52.6		10449	10431	50.9	47.4	0.5	520	75.4	. 41	28	53.2
	1053	9930	9948	51.5	52.6		10449	10428	51.9	40.9	0.4	520	75.4	41	28	53.3
	1054	9929	9948	53.2	50		10449	10425	54.6	40	1.4	521	75.4	40.9	28	53.8
35	1055	9931	9952	53	45.5		10449	10425	54.6	40	1.6	519	75.3	40.8	28	53.7
	1056	3791	3808	50	50		4445	4425	50.6	42.9	0.5	655	75.5	40.5	28	52.9
37	1057	3791	3808	50	50		4444	4424	50.6	42.9	0.5	654	75.5	40.5	28	53
	1058	9930	9949	52.2		3558	10449	10431	50.9	47.4	1.2	520	75.4	41	28	53.2
	1059	9930	9949	52.2		3559	10449	10428	51.9	40.9	0.3	520	75.4	41	28	53.5
	1060	3788	3805	50		3560	4445	4425	50.6	42.9	0.5	658	75.5	40.4	28	52.9
	1061	19800	19817	50.4		3561	19921	19900	51.8	45.5	1.4	122	72.2	43.4	28	50.8
	1062	3787	3804	50		3562	4444	4424	50.6	42.9	0.5	658	75.5	40.4	28	52.9
	1063	25782	25806	53.5		3563	26183	26163	51.7	42.9	1.7	402	74.7	40.3	28	52.9
	1064	25782	25806	53.5		3564	26183	26160	54.5	41.7	1	402	74.7	40.3	28	53.5
	1065	25782	25806	53.5		3565	26183	26159	54.9	40	1.5	402	74.7	40.3	28	53.5
	1066	2429	2447	50.2		3566	3187	3166	50.3	45.5	0.1	759	76.6	43	29	53.8
	1067	2427	2445	52.1		3567	3185	3164	51	45.5	1.1	759	76.7	. 43.1	29	54.1
	1068	2429	2447	50.2		3568	3185	3164	51	45.5	0.7	757	76.6	42.9	29	53.8
	1069	19800	19817	50.4		3569	19923	19904	50.1	50	0.3	124	72.3	43.5	29	50.8
	1070	2427	2445	52.1		3570	3187	3166	50.3	45.5	1.8	761	76.7	43.1	29	53.9
	1071	29183	29204	50.4		3571	29412	29393	50.3	45	0	230	75.3	44.8	29	52.9
	1072	16367	16386	51.4		3572	16780	16760	51.4	42.9	0.1	414	75	40.8	30	53
	1073	11543	11562	50.4	40	3573	12254	12236	50.5	47.4	0.1	712	76.2	42	30	53.6
	1074	12976	12995	51.1	45	3574	13547	13528	50.2	45	0.9	572	77.4	45.5	30	54.3
	1075	12040	12057	50.6		3575	12254	12236	50.5	47.4	0.1	215	75.5	45.6	30	53.1
	1076	12976	12996	51.8	42.9	3576	13544	13525	52.6	55	0.8	569	77.5	45.7	30	54.8
	1077	10141	10160	51	45	3577	10605	10588	51.1	50	0.1	465	74.9	40.2	30	52.8
	1078	12235	12253	50.1		3578	12996	12977	50.2	40	0.1	762	76.4	42.4	30	53.6
	1079	19795	19814	50.4		3579	19921	19901	50.2	47.6	0.3	127	72.3	43.3	30	50.8
	1080	12235	12253	50.1	52.6		12993	12975	51.4	47.4	1.3	759	76.5	42.6	30	53.7
	1081	12976	12994	50.3	47.4	3581	13547	13528	50.2	45	0.1	572	77.4	45.5	30	54.3
	1082	12975	12994	52.1		3582	13544	13525	52.6	55	0.5	570	77.4	45.6	30	54.9
	083	12977	12996	50.2		3583	13547	13528	50.2	45	0	571	77.3	45.4	30	54.3
	084	11541	11561	50.9		3584	12254	12236	50.5	47.4	0.3	714	76.2	42	30	53.6
65 1	1085	28394	28411	50.3	50	3585	28672	28654	50.6	52.6	0.3	279	78.6	51.6	30	55.2
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Gel 1086 9930 9948 51.5 52.6 3586 10455 10495 51.4 40.9 0.3 522 75.3 40.7 30 53.1																
68 1088 9990 9940 52.2 50 3588 10455 10435 50.5 42.9 1.7 528 75.3 40.7 30 52.2 68 1089 12236 12256 51.2 42.9 9389 12412 12392 50 42.9 1.2 177 73 41.2 30 51.2 77 1091 9930 9949 52.2 50 3580 10455 10435 50.5 42.9 0.4 523 75.2 40.5 30 52.9 77 1092 12726 12746 51.3 47.6 5389 10455 10435 50.5 42.9 0.4 523 75.2 40.5 30 52.9 78 1093 9933 9952 50.9 45 3593 10455 10435 50.5 42.9 0.4 523 75.2 40.5 30 52.9 79 1093 9933 9952 50.9 45 3593 10455 10434 51.1 40.9 0.3 523 75.2 40.5 30 52.9 79 1094 16909 16928 50.8 45 3593 10455 10434 51.1 40.9 0.3 523 75.2 40.5 30 52.9 76 1096 12976 12993 51.4 47.4 3598 1367 3168 50.3 45.5 50 0.3 52.7 77.7 45.5 30 52.7 76 1098 2677 2692 52.1 40.9 3599 3167 3168 50.3 45.5 1.8 517 75.6 41.6 30 53.1 77 1097 19800 19816 52.1 52.6 5399 10455 10435 50.5 42.9 1 526 75.3 40.7 30 52.9 78 1098 12375 12393 51.4 47.4 3598 13647 13628 50.2 45 1.2 573 77.3 45.4 30 54.3 79 1098 9930 9948 51.5 52.6 3599 10455 10435 50.5 42.9 1 526 75.3 40.7 30 52.9 80 1100 12976 12995 51.1 45.3 300 13644 13525 52.6 55 1.5 560 77.5 45.4 30 54.3 80 1101 24635 24635 50.5 52.6 3602 5182 25164 51.4 47.4 0.9 548 75.1 40.7 30 52.8 80 1103 24630 24641 50.5 52.6 3602 25182 25164 51.4 47.4 0.9 548 75.1 40.7 30 52.8 80 1103 24630 24641 51.5 42.9 3609 26569 26571 51.7 47.4 0.9 548 75.1 40.1 30 52.8 80 1103 24630 24641 51.5 42.9 3609 26569 26571 51.7 47.4 0.9 548 75.1 40.1 30 52.8 80 1101 24635 26647 50.1 52.6 3613 26680														40.7		
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To 1090					52.2			10435	50.5		1.7	'526	75.3	40.7	30	52.9
Tr	69	1089	12236	12256	51.2	42.9 3589	12412	12392	50	42.9	1.2	177	73	41.2	30	51.2
Transfer	70	1090	9930	9949	52.2	50 3590	10455	10434	51.1	40.9	1.1	.526	75.3	40.7	30	53.1
T3 1093	71	1091	9933	9952	50.9	45 3591	10455	10435	50.5	42.9	0.4	523	75.2	40.5	30	52.9
Train 1994 16999 16928 50.8 45 3594 17501 17461 51.2 42.9 0.4 559 75.9 41.8 30 53.5 75 10957 12995 51.4 47.4 3595 13544 13525 52.6 55 1.2 570 77.4 45.6 30 54.7 76 1096 2671 2692 52.1 40.9 3596 3167 3166 50.3 45.5 1.8 511 75.6 41.6 30 53.7 77 1097 10900 19816 52.1 52.6 3597 10921 10900 51.8 45.5 0.3 122 72.2 43.4 30 51.2 78 1009 12975 12995 51.4 47.4 3598 13547 13528 50.2 45 1.2 573 77.3 45.4 30 51.2 51.0 30	72	1092	12726	12746	51.3	47.6 3592	13314	13297	51	50	0.3	'589	76.6	43.6	30	54
To 1995 12975 12993 51.4 47.4 3595 13544 13525 52.6 55 1.2 570 77.4 45.6 30 54.7	73	1093	9933		50.9			10434	51.1	40.9	0.3	523	75.2	40.5	30	
To 1096	74	1094	16909	16928	50.8	45 3594	17501	17481	51.2	42.9	0.4	593	75.9	41.8	30	53.5
Tr 1997 19800 19818 52.1 52.6 3597 19921 19900 51.8 45.5 0.3 122 72.2 43.4 30 51.2 78 1098 12975 12993 51.4 47.4 3598 13547 13528 50.2 45 1.2 573 77.3 45.4 30 63.2 63 79 1099 9900 9900 9946 51.5 52.6 3590 10455 10435 50.5 42.9 1 52.6 75.3 40.7 30 52.9 60 1100 12976 12995 51.1 45 3600 13544 13525 52.6 55 1.5 589 77.5 45.7 30 54.6 61 1101 24635 24653 50.5 52.6 3601 25182 25184 51.4 47.4 0.9 548 75.1 40.1 30 52.6 3602 25182 25184 51.4 47.4 1.3 5507 52.2 40.2 30 52.7 63 1103 24630 24648 50.8 52.6 3603 25182 25184 51.4 47.4 1.3 5507 52.2 40.2 30 52.7 63 1103 24630 24648 50.8 52.6 3603 25182 25184 51.4 47.4 1.3 5507 52.2 40.3 30 53 34 1104 28394 28412 51.1 47.4 3004 28672 28684 50.6 52.6 0.6 277 78.6 51.6 30 55.3 37 1107 28421 28413 50.2 241 3605 28672 28684 50.6 52.6 0.6 277 78.6 51.6 30 55.3 37 1107 28421 28441 51.5 42.9 3607 26587 26588 52.7 45 1.2 167 72.3 40.1 30 51.2 80 1109 28421 28441 51.5 42.9 3609 26589 26572 51 50 0.5 169 72.4 40.2 30 51.1 30 1110 26421 26441 51.5 42.9 3609 26589 26572 51 50 0.5 169 72.4 40.2 30 51.3 30 3113 20309 26057 52.6 52.6 3612 26183 26183 51.7 50 0.3 170 72.3 40.1 30 51.2 51 51 51 51 51 51 51 5	75	1095	12975	12993	51.4	47.4 3595	13544	13525	52.6	55	1.2	570	77.4	45.6	30	54.7
T8 1098 12975 12993 51.4 47.4 3598 13547 13528 50.2 45 1.2 573 77.3 45.4 30 54.3 T9 1099 9900 9948 51.5 52.6 3599 10455 10435 50.5 42.9 1 526 75.3 40.7 30 54.6 S0 1100 12976 12995 51.1 45 3600 13544 13525 52.6 55 1.5 569 77.5 45.7 30 54.6 S1 1101 24635 24653 50.5 52.6 3601 25182 25184 51.4 47.4 0.9 548 75.1 40.1 30 52.8 S2 1102 24633 24651 50.1 52.6 3602 25182 25184 51.4 47.4 0.9 548 75.1 40.1 30 52.8 S3 1103 24630 24648 50.8 52.6 3603 25182 25184 51.4 47.4 0.9 548 75.1 40.1 30 52.8 S3 1103 24630 24648 50.8 52.6 3603 25182 25184 51.4 47.4 0.9 548 75.1 40.1 30 52.8 S4 1104 28394 28412 51.1 47.4 3604 28672 28654 50.6 52.6 0.5 279 78.6 51.6 30 55.3 S5 1105 28395 28413 50.2 42.1 3605 28672 28654 50.6 52.6 0.6 277 78.6 51.6 30 55.3 S7 1107 24221 26441 51.5 42.9 3607 26587 26568 52.7 45 1.2 167 72.3 40.1 30 51.2 S6 1108 28421 26441 51.5 42.9 3607 26589 26572 51 50 0.5 169 72.4 40.2 30 51.1 S9 1110 26421 26441 51.5 42.9 3609 26589 26573 51.7 50 0.3 170 72.3 40.1 30 51.2 S7 1111 26040 26061 56.4 54.5 3611 26599 26572 51 50 0.5 169 72.4 40.2 30 51.2 S9 1111 26039 26057 52.6 52.6 3613 26182 26183 51.7 42.9 0.9 14.6 14.7 40.7 30 51.2 S8 1118 11541 11560 50.1 45 3616 26183 26160 54.5 41.7 1.9 445 71.9 40.7 30 51.2 S9 1111 11640 11557 50.4 50 3617 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 S9 1111 11540 11567 50.4 50 3617 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 S9 1111 11540 11567 50.4 50 3617 1	76	1096	2671	2692	52.1	40.9 3596	3187	3166	50.3	45.5	1.8	517	75.6	41.6	30	53.1
No. 1099 9930 9948 51.5 52.6 3599 10455 10435 50.5 42.9 1 526 75.3 40.7 30 52.9	77	1097	19800	19818	52.1	52.6 3597	19921	19900	51.8	45.5	0.3	122	72.2	43.4	30	51.2
B0 1100 12976 12995 51.1 45 3600 13544 13525 52.6 55 1.5 580 77.5 45.7 30 54.6 B1 1101 24635 24683 50.5 52.6 3601 25182 25184 51.4 47.4 0.9 54.8 75.1 40.1 30 52.5 B3 1103 24630 24648 50.6 52.6 3603 25182 25184 51.4 47.4 1.3 5505 75.2 40.2 30 52.7 B3 1103 24630 24648 50.8 52.6 3603 25182 25184 51.4 47.4 0.6 553 75.2 40.3 30 53 B4 1104 28394 28412 51.1 47.4 39.04 26672 28654 50.6 52.6 0.4 279 78.6 51.6 30 53.3 B5 1105 28395 28413 50.2 42.1 3605 28672 28654 50.6 52.6 0.4 278 78.8 51.4 30 55.2 B6 1106 28395 28415 51.2 45 3606 28672 28654 50.6 52.6 0.4 278 78.8 51.4 30 55.2 B6 1106 28395 28415 51.2 45 3606 28672 28654 50.6 52.6 0.6 277 78.6 51.6 30 55.3 B7 1107 26421 26441 51.5 42.9 3609 26589 26571 51.7 47.4 0.2 169 72.4 40.2 30 51.2 B8 1108 26421 26441 51.5 42.9 3609 26589 26571 51.7 47.4 0.2 169 72.4 40.2 30 51.2 B9 1110 26421 26441 51.5 42.9 3609 26589 26572 51 50 0.5 169 72.4 40.2 30 51.2 B9 1111 26040 26061 564 64.5 3611 26589 26589 55.2 45.5 1.2 550 75.1 40 30 54.2 B9 1111 26040 26061 564 64.5 3611 26589 26589 55.2 45.5 1.2 550 75.1 40 30 54.2 B9 1111 26039 26057 52.6 52.6 3614 26183 26163 51.7 50 0.3 170 72.3 40.3 30 50.7 B9 1115 8867 8887 52.3 47.6 3618 10605 10588 51.1 50 0.6 359 74.6 40.4 30 52.4 50 50 51.1 50 50 51.1 50 50.5 51.1 50 50.5 51.1 50 50.5 51.1 50 50.5 51.1 50.0 51.1 50.0 51.2 50.5 51.1 50.0 51.2 50.5 51.1 50.0 51.2 50.5 51.1 50.0 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5 50.5	78	1098	12975	12993	51.4	47.4 3598	13547	13528	50.2	45	1.2	573	77.3	45.4	30	54.3
81 1101	79	1099	9930	9948	51.5	52.6 3599	10455	10435	50.5	42.9	1	526	75.3	40.7	30	52.9
82 1102 24833 24851 50.1 52.6 3602 25182 25164 51.4 47.4 1.3 550 75.2 40.2 30 52.7 83 1103 24630 24648 50.8 52.6 3603 25182 25164 51.4 47.4 0.6 553 75.2 40.3 30 55.3 85 1105 28395 28413 50.2 42.1 3605 28672 28654 50.6 52.6 0.4 278 78.6 51.4 30 55.2 86 1106 28395 28413 50.2 42.1 453 800 28672 28654 50.6 52.6 0.6 277 78.6 51.6 30 55.2 4861 21 267.23 40.1 2861 2861 2861 28.6 0.6 22.7 78.6 51.6 30 55.2 36.1 30 55.2 36.1 30 36.1 30 35.1 39 28.1 21.7 22.3	80	1100	12976	12995	51.1	45 3600	13544	13525	52.6	55	1.5	;569	77,5	45.7	30	54.6
83 1103	81	1101	24635	24653	50.5	52.6 360	25182	25164	51.4	47.4	0.9	-548	75.1	40.1	30	52.8
84 1104 28394 28412 51.1 47.4 3604 28672 28654 50.6 52.6 0.5 279 78.6 51.6 30 55.3 85 1105 28395 28413 50.2 42.1 3605 28672 28654 50.6 52.6 0.4 278 78.6 51.6 30 55.3 86 1106 28396 28415 51.2 45 3606 28672 28654 50.6 52.6 0.6 277 78.6 51.6 30 55.3 87 1107 28421 28441 51.5 42.9 3607 26587 26586 52.7 45 1.2 167 72.3 40.1 30 51.2 88 1108 28421 28441 51.5 42.9 3607 26587 26586 52.7 45 1.2 167 72.3 40.1 30 51.2 89 1109 28421 28441 51.5 42.9 3609 26599 26571 51.7 47.4 0.2 169 72.4 40.2 30 51.1 90 1110 28421 28441 51.5 42.9 3610 26599 26572 51 50 0.5 169 72.4 40.2 30 51.1 91 1111 28040 28061 56.4 54.5 3611 28599 26589 26572 51 50 0.5 169 72.4 40.2 30 51.2 92 1112 26039 26057 52.6 52.6 3613 26180 26180 54.5 41.7 1.9 145 71.9 40.7 30 51.2 93 1113 26039 26057 52.6 52.6 3613 26182 26181 51.2 40.9 0.9 1.4 144 71.7 40.3 30 50.7 94 1114 26039 26057 52.6 52.6 52.6 3614 26183 26180 54.5 41.7 1.9 145 71.9 40.7 30 51.2 95 1116 10247 10267 50.5 47.6 3616 10605 10588 51.1 50 0.6 39 74.6 40.4 30 52.4 97 1117 11540 11557 50.4 50 5618 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 50 11.1 11541 11560 50.1 45 5618 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 50 11.1 11541 11560 50.1 45 5618 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 50 11.1 11541 11560 50.1 45 5618 12254 12236 50.5 47.4 0.4 714 76.2 42.1 30 53.6 50 50 50 50 50 50 50 5	82	1102	24633	24651	50.1	52.6 3602	25182	25164	51.4	47.4	1.3	,550	75.2	40.2	30	52.7
S 1105	83	1103	24630	24648	50.8	52.6 3603	25182	25164	51.4	47.4	0.6	553	75.2	40.3	30	53
Ref 1106 28396 28415 51.2 45 3606 28672 28654 50.6 52.6 0.6 277 78.6 51.6 30 55.3 Ref 1107 28421 28441 51.5 42.9 3607 28587 28588 28.7 45 1.2 167 72.3 40.1 30 51.2 Ref 1808 1108 28421 28441 51.5 42.9 3608 28699 28571 51.7 47.4 0.2 169 72.4 40.2 30 51.2 Ref 1109 28421 28441 51.5 42.9 3608 28589 28571 51.7 50 0.5 169 72.4 40.2 30 51.1 Ref 1110 28421 28441 51.5 42.9 3608 28589 28572 51 50 0.5 169 72.4 40.2 30 51.1 Ref 1111 28040 28041 51.5 42.9 3608 28589 28587 51.7 50 0.3 170 72.3 40 30 51.2 Ref 1111 28040 28061 584 45.5 3611 28589 28586 55.2 45.5 1.2 550 75.1 40 30 54.2 Ref 1112 28039 28057 52.6 52.6 3612 26183 26180 54.5 41.7 1.9 146 71.9 40.7 30 51.2 Ref 1112 28039 28057 52.6 52.6 3612 26183 26180 54.5 41.7 1.9 146 71.9 40.7 30 51.2 Ref 1116 28639 28657 52.6 52.6 3614 26183 26183 51.7 40.9 1.4 144 71.7 40.3 30 50.2 Ref 1116 10247 10267 50.5 47.6 3616 10268 51.6 51.2 40.9 1.4 144 71.9 40.7 30 51.2 Ref 1116 10247 10267 50.5 47.6 3616 10268 51.6 51.2 40.9 1.4 144 71.9 40.7 30 51.2 Ref 1116 10247 10267 50.5 47.6 3616 10268 51.6 51.5 50 6.8 39 74.6 40.4 30 53.4 Ref 1116 11267	84	1104	28394	28412	51.1	47.4 3604	28672	28654	50.6	52.6	0.5	279	78.6	51.6	30	55.3
87 1107 26421 26441 51.5 42.9 3607 26587 26568 52.7 45 1.2 167 72.3 40.1 30 51.2 88 1108 26421 26441 51.5 42.9 3608 26598 26571 51.7 47.4 0.2 169 72.4 40.2 30 51.1 99 1109 26421 26441 51.5 42.9 3608 26589 26572 51 50 0.5 169 72.4 40.2 30 51.1 90 1110 26421 26441 51.5 42.9 3610 26599 26572 51 50 0.5 169 72.4 40.2 30 51.1 91 1111 26040 26061 56.4 54.5 3611 25599 25658 52.7 55 0.3 170 72.3 40 30 51.2 91 1111 26039 26057 52.6 52.6 3612 26183 26160 54.5 41.7 1.9 41.7 40.3 30 50.7 93 1113 26039 26057 52.6 52.6 3613 26182 26181 51.2 40.9 1.4 444 71.7 40.3 30 50.7 94 1114 26039 26057 52.6 52.6 3613 26182 26181 51.2 40.9 1.4 144 71.7 40.3 30 50.7 94 1114 26039 26057 52.6 52.6 3614 26183 26183 51.7 42.9 0.9 145 71.9 40.7 30 51.2 95 1115 8867 8887 52.3 47.6 3616 29253 9235 51.6 47.4 0.7 367 57.5 41.3 30 53.2 96 1116 10247 10267 50.5 47.6 3616 10605 10588 51.1 50 0.6 359 74.6 40.4 30 52.4 97 1117 11540 11557 50.4 50 3618 12254 12236 50.5 47.4 0.4 714 76.2 42.1 30 53.6 98 1118 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.4 714 76.2 42.1 30 53.6 100 1120 13039 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 46 31 52.2 102 1122 19709 19730 51.3 40.9 3622 19921 19900 51.8 45.5 0.5 213 73.9 41.8 31 52.2 105 1125 4 2 2 52.3 52.6 3624 256 235 52.6 45.5 0.8 217 71.9 71.6 40.1 30 53.6 107 1127 13900 13915 51.8 50 3628 13177 13156 50.4 40.9 0.4 70.9 76.6 43.2 31 54.2 107 1127 13900 13915 51.8 50 3628 13177	85	1105	28395	28413	50.2	42.1 3605	28672	28654	50.6	52.6	0.4	278	78.6	51.4	30	55.2
B8 1108 26421 26441 51.5 42.9 3608 26589 26571 51.7 47.4 0.2 169 72.4 40.2 30 51.2 B8 1109 26421 26441 51.5 42.9 3600 26589 26572 51 50 0.5 169 72.4 40.2 30 51.2 S9 1110 26421 26441 51.5 42.9 3600 26589 26573 51.7 50 0.3 170 72.3 40 30 51.2 S9 1111 26040 26061 56.4 54.5 3611 26589 26568 55.2 45.5 1.2 550 75.1 40 30 54.2 S9 1112 26039 26057 52.6 52.6 3612 26182 26160 54.5 41.7 1.9 145 71.9 40.7 30 54.2 S9 1113 26039 26057 52.6 52.6 3613 26182 26161 51.2 40.9 1.4 144 71.7 40.3 30 50.7 S9 1114 26039 26057 52.6 52.6 3614 26183 26163 51.7 42.9 0.9 145 71.9 40.7 30 51.2 S9 1115 8867 8867 82.3 47.6 3615 2653 2258 51.6 47.4 0.7 367 75.1 41.3 30 52.4 S9 1116 10247 10267 50.5 47.6 3616 10605 10588 51.1 50 66 367 48.6 40.4 30 52.4 S9 1118 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 S9 1119 8221 8240 524 50 3619 8929 8911 53.4 52.6 1.7 76.2 42.1 30 53.6 S9 1119 1921 19309 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 46 31 52 S1 101 1121 13901 13919 53.2 52.6 3624 2563 3621 19917 13895 50.5 47.4 0.1 71.9 75.4 40.1 31 53.8 S1 104 1124 3 21 53.4 52.6 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.8 S1 106 1126 13039 13057 51.1 52.6 3623 16774 16751 53.6 41.7 0.8 40.9 75.1 41.1 31 53.8 S1 107 1127 13900 13917 50.4 50.3628 13177 13156 50.4 40.9 0.7 139 73.9 41.8 13 52.2 S1 101 1121 13901 13918 53.2 52.6 3623 16774 16751 53.6 41.7 0.8 40.9 75.1 41.1 31 53.8 S1 101 11	86	1106	28396	28415	51.2	45 3606	28672	28654	50.6	52.6	0.6	'277	78.6	51.6	30	55.3
89 1109 26421 26441 51.5 42.9 3609 26589 26572 51 50 0.5 169 72.4 40.2 30 51.1	87	1107	26421	26441	51.5	42.9 3607	26587	26568	52.7	45	1.2	167	72.3	40.1	30	:51.2
90 1110 26421 26441 51.5 42.9 3610 26590 26573 51.7 50 0.3 170 72.3 40 30 51.2 91 1111 26040 26061 56.4 54.5 3611 26599 26568 55.2 45.5 1.2 550 75.1 40 30 54.2 92 1112 26039 26057 52.6 52.6 3612 26183 26160 54.5 41.7 1.9 14.7 40.3 30 54.2 93 1113 26039 26057 52.6 52.6 3613 26182 26160 54.5 41.7 1.9 14.7 40.3 30 50.7 94 1114 26039 26057 52.6 52.6 3614 26183 26160 54.5 41.7 1.9 14.7 40.3 30 50.7 94 1114 26039 26057 52.6 52.6 3614 26183 26160 54.5 41.7 1.9 14.7 40.7 30 51.2 95 1115 8867 8867 52.3 47.6 3615 9253 9235 51.6 47.4 0.7 367 75.1 41.3 30 53.2 96 1116 10247 10267 50.5 47.6 3615 10605 10588 51.1 50 0.6 359 74.6 40.4 30 52.4 97 1117 11540 11557 50.4 50 3618 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 98 1118 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.4 714 76.2 42 30 53.6 100 1120 13039 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 41.8 31 52.2 102 1122 19709 19730 51.3 40.9 3622 19921 19900 51.8 45.5 0.5 17.3 73.9 41.8 31 52.2 105 1125 42 22 52.3 52.6 3626 13177 13156 50.4 40.9 0.7 139 73.9 41.8 31 52.2 106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 0.2 117 71.7 71.8 40.1 31 53.8 106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 0.2 117 71.7 71.8 40.1 31 53.8 107 1127 13900 13917 50.4 50.8624 256 205 50.2 40.5 0.8 254 76.1 40.1 31 53.8 107 1127 13900 13957 51.1 50.8 3629 13177 13156 50.4 40.9 0.4 70.9 76.6 43.2 31 54.1 109 1129 13039 13057 51.1 52.6 3629 13177 13	88	1108	26421	26441	51.5	42.9 3608	26589	26571	51.7	47.4	0.2	169	72.4	40.2	30	51.2
11111 26040 26061 56.4 54.5 3611 26599 26568 55.2 45.5 1.2 550 75.1 40 30 54.2	89	1109	26421	26441	51.5	42.9 3609	26589	26572	51	50	0.5	169	72.4	40.2	30	51.1
92 1112 26039 26057 52.6 52.6 3612 26183 26160 54.5 41.7 1.9 145 71.9 40.7 30 51.2 39 1113 26039 26057 52.6 52.6 3613 26182 26163 51.2 40.9 1.4 144 71.7 40.3 30 50 96 1115 8867 8887 52.3 47.6 3615 9253 9235 51.6 47.4 0.7 367 75.1 41.3 30 53.2 96 1116 10247 10267 50.5 47.6 3616 10505 15.6 47.4 0.7 367 75.1 41.3 30 53.2 96 1116 10247 10267 50.5 47.6 3616 10605 10.1 11.7 17.7 75.1 41.3 30 53.2 98 1118 11541 11567 50.4 50.3 81.7 <td>90</td> <td>1110</td> <td>26421</td> <td>26441</td> <td>51.5</td> <td>42.9 3610</td> <td>26590</td> <td>26573</td> <td>51.7</td> <td>50</td> <td>0.3</td> <td>170</td> <td>72.3</td> <td>. 40</td> <td>30</td> <td>51.2</td>	90	1110	26421	26441	51.5	42.9 3610	26590	26573	51.7	50	0.3	170	72.3	. 40	30	51.2
93 1113 26039 26057 52.6 52.6 3613 26182 26161 51.2 40.9 1.4 144 71.7 40.3 30 50.7 41114 26039 26057 52.6 52.6 3614 26183 26163 51.7 42.9 0.9 145 71.9 40.7 30 51.7 51.1 41.3 30 53.7 51.1 41.3 41.3 41.3 41.3 41.3 41.3 41.3 4	91	1111	26040	26061	56.4	54.5 3611	26589	26568	55.2	45.5	1.2	550	75.1	40	30	54.2
94 1114 26039 26057 52.6 52.6 3614 26183 26163 51.7 42.9 0.9 145 71.9 40.7 30 51 .95 1115 8867 887 52.3 47.6 3615 9255 9256 51.6 47.4 0.7 307 75.1 41.3 30 52.5 .96 1116 10247 10267 50.5 47.6 3616 10605 10588 51.1 50 .66 359 47.8 40.4 0.3 05.2 4.9 1117 11540 11557 50.4 50 3617 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 98 11.8 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 98 11.8 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.4 714 76.2 42.1 30 53.6 99 1119 0821 8240 524 52 40 5619 8929 8911 53.4 52.6 1 7.09 75.4 40.1 30 53.8 100 1120 13039 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 46 31 52 101 1121 13901 13919 53.2 52.6 3621 13917 13956 50.4 40.9 0.7 139 73.9 46 31 52 101 1121 13901 13919 53.2 52.6 3621 13917 13956 50.4 40.9 0.7 139 73.9 46 31 52 102 1122 19709 19730 51.3 40.9 3622 19921 19900 51.8 45.5 0.5 217.3 94 11.8 13 52.8 104 1124 3 21 53.4 52.6 3624 256 25 52.6 36.4 40.9 15.8 50.5 47.4 1.7 1.8 40.6 13.1 52.8 106 1126 13039 13058 51.8 52.6 3624 256 25 52.6 36.4 40.9 1.5 13.9 73.9 46 31 52.2 105 1125 4 22 52.3 52.6 3624 256 13177 13156 50.4 40.9 15.5 13.9 73.9 46 31 52.2 105 1125 4 22 52.3 52.6 3624 256 25 52.6 36.4 40.9 15.5 13.9 14.1 31 53.8 104 1124 3 21 53.4 52.6 3624 256 25 52.6 36.5 0.5 217.6 17.6 84.6 13 53.8 104 1124 3 22 52.3 52.6 3625 314 296 50.0 47.4 1.7 317.6 84.6 13 54.1 106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.2 107 1127 13900 13917 50.4 50.3 50.8 1377 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 109 1129 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 109 1129 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 109 1129 13039 13058 51.8 50 3628 13774 13726 50.8 40.9 0.4 70.9 76.6 43.2 31 54.1 109 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.4 70.9 76.6 43.2 31 54.1 111 131 3 24148 24436 50 24.4 9.3 633 24527 24508 50.5 40.9 0.4 70.9 76.6 43.2 31 54.1 111 131 3 24158 24305 50 47.6 3633 24527 24508 50.5 50.5 40.9 0.2 107 71.7 14.2 71 150.5 11.5 13.6 13.5 13.5 13.	92	1112	26039	26057	52.6	52.6 3612	26183	26160	54.5	41.7	1.9	145	71.9	40.7	30	51.2
No.	93	1113	26039	26057	52.6	52.6 3613	26182	26161	51.2	40.9	1.4	144	71.7	40.3	30	50.7
116	94	1114	26039	26057	52.6	52.6 3614	26183	26163	51.7	42.9	0.9	145	71.9	· 40.7	30	51
97 1117 11540 11557 50.4 50 3617 12254 12236 50.5 47.4 0.1 715 76.2 42.1 30 53.6 98 1118 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.4 714 76.2 42.1 30 53.6 98 1119 6221 8240 524 524 50 3619 8929 8911 53.4 52.6 1 7.09 75.4 40.1 30 53.6 100 1120 13039 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 46 31 52 101 1121 13901 13919 53.2 52.6 3621 13917 13956 50.4 40.9 0.7 139 73.9 46 31 52 102 1122 19709 19730 513.4 40.9 3622 19921 19900 51.8 45.5 0.5 131 73.9 41.8 13 52.2 102 1122 19709 19730 513.4 40.9 3622 19921 19900 51.8 45.5 0.5 131 73.9 41.8 13 52.2 102 1122 19709 19730 513.4 40.9 3622 19921 19900 51.8 45.5 0.5 131 73.9 41.8 13 52.2 103 1123 16366 16386 54.4 52.4 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.8 104 1124 3 21 53.4 52.6 3624 256 25 52.6 35.6 45.5 0.8 254 76.1 46.1 31 52.2 105 1125 4 22 52.3 52.6 3624 256 25 52.6 45.5 0.8 254 76.1 46.1 31 54.2 106 1126 13039 13058 51.8 50 3628 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 106 1126 13039 13058 51.8 50 3628 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 107 1127 13900 13917 50.4 50 3628 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 109 1129 13039 13058 51.8 50 3628 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 109 1129 13039 13058 51.8 50 3628 13177 1376 50.8 40.9 0.2 117 71.7 42.7 31 50.1 109 1129 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.2 117 71.7 42.7 31 50.1 111 131 3 21 53.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 111 131 3 24418 24436 50 24 47.6 3632 27444 53 42.9 50.8 44.9 0.2 100 70.8 43 31 50.6 113 1133 24418 24430 50 47.4 3633 24527 24508 50.5 40.9 0.2 100 70.8 43 31 50.6 113 1133 24418 24430 50 47.4 3633 24527 24508 50.5 44.4 0 756 75.9 41.1 31 53.1 151 135 24719 24200 53.3 40.9 3635 24936 24919 51.8 50 44.4 0 756 75.9 41.1 31 53.7 115 1135 24719 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 411 31 53.7	. 95	1115	8867	8887	52.3	47.6 3618	9253	9235	51.6	47.4	0.7	387	75.1	41.3	30	53.2
98 1118 11541 11560 50.1 45 3618 12254 12236 50.5 47.4 0.4 714 76.2 42 30 53.5 99 1119 8221 8240 52.4 503 3619 8929 8911 53.4 52.6 1 709 75.4 40.1 30 53.5 100 1120 13039 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 137.9 46 31 52 101 1121 19801 19819 53.2 52.6 3621 19917 19895 52.5 43.5 0.8 117 72 43.6 31 51.2 102 1122 19709 19730 51.3 40.9 3622 19917 19895 52.5 43.5 0.8 117 72 43.6 31 51.2 103 1123 16366 16386 54.4 52.4 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.8 104 1124 3 21 53.4 52.6 3624 256 255 52.6 45.5 0.8 254 76.1 46.1 31 54.2 105 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13056 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52.1 107 1127 19800 19817 50.4 50 3627 19916 19895 50.2 40.9 0.2 117 71.7 42.7 31 50.3 108 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50 0.5 662 75.6 40.8 31 53.1 110 1129 13039 13056 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 110 1130 13039 13056 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 111 1131 3 2416 24336 50 47.4 3633 24527 24508 50.5 40.9 0.2 100 70.8 43.2 31 54.1 111 1131 24438 50 47.4 3633 24527 24508 50.5 44.4 0 756 75.9 41.1 31 53.7 116 1135 24478 24430 50 47.4 3633 24527 24508 50.5 44.4 0 756 75.9 41.1 31 53.7 116 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 4.4, 0 756 75.9 41.1 31 53.7 116 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 4.4, 0 756 75.9 41.1 31 53.7	96	1116	10247	10267	50.5	47.6 3616	10605	10588	51.1	50	0.6	359	74.6	40.4	30	52.4
99 1119 8221 8240 52.4 50 3619 8929 8911 53.4 52.6 1 709 75.4 40.1 30 53.6 100 1120 19399 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 46 31 51.2 101 1121 19801 19819 53.2 52.6 3621 19917 19895 52.5 34.5 0.8 117 72 43.6 31 51.2 102 1122 19709 19730 51.3 40.9 3622 19921 19900 51.8 45.5 0.5 213 73.9 41.8 31 52.2 103 1123 16366 16366 63.4 52.4 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.8 104 1124 3 21 53.4 52.6 3625 3625 235 52.6 45.5 0.8 25.4 76.1 46.1 31 53.2 105 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13058 51.8 50 3626 13177 13166 50.4 40.9 1.5 139 73.9 46 31 52.1 107 1127 19800 19817 50.4 500 3627 19916 19895 50.2 40.9 0.2 117 71.7 42.7 31 50.3 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54.1 112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 13 1133 24418 24436 50 47.4 3633 24527 24508 50.5 44.4 0 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8	97	1117	11540	11557	50.4	50 3617	12254	12236	50.5	47.4	0.1	715	76.2	42.1	30	153.6
1100 1120 13039 13057 51.1 52.6 3620 13177 13156 50.4 40.9 0.7 139 73.9 46 31 52 101 1121 13901 13919 53.2 52.6 3621 13917 13935 52.5 43.5 0.8 117 72 43.6 31 51 52 102 1122 13709 13935 53.4 40.9 3622 13921 13990 51.8 45.5 0.5 213 73.9 41.8 31 52.2 103 1123 16366 16386 54.4 52.4 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.8 104 1124 3 21 53.4 52.6 3624 256 225 235 6.6 45.5 0.8 254 76.1 48.1 31 53.8 104 1124 3 21 53.4 52.6 3624 256 235 52.6 45.5 0.8 254 76.1 48.1 31 54.2 105 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52 107 1127 13900 13917 50.4 50 3627 13916 13985 50.2 40.9 0.2 117 71.7 42.7 31 53.8 1109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 4311 131 3 21 53.4 50 3630 27464 27444 53 42.9 0.2 100 70.8 43.3 31 50.6 113 133 24418 24436 50 47.4 3633 24527 24508 50.5 50.5 40.9 0.2 107 70.8 43.3 31 50.6 113 1133 24418 24436 50 47.4 3633 24527 24508 50.5 50.5 40.7 75.8 75.8 41.1 31 53.7 116 1135 24708 26727 50.4 43.9 3634 27463 27446 50 44.4 0.7 75.8 75.9 41.1 31 53.7 116 1135 24719 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 75.8 75.8 41.1 31 53.7 116 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 75.8 75.8 41.1 31 53.7 116 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 75.8 75.8 41.1 31 53.7 116 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 75.8	98	1118	11541	11560	50.1	45 3618	12254	12236	50.5	47.4	0.4	714	76.2	42	30	53.5
101 1121 19801 19819 53.2 52.6 3621 19917 19895 52.5 43.5 0.8 '117 72 43.6 31 51.2 1902 1122 19709 19730 51.3 40.9 3622 19921 19900 51.8 45.5 0.5 213 73.9 41.8 31 52.8 103 113 16368 54.4 52.4 3623 16774 16751 53.6 41.7 0.8 40.9 75.1 41.1 31 53.8 104 1124 3 21 53.4 52.6 3624 256 235 52.6 45.5 0.8 254 76.1 46.1 31 54.2 105 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13058 51.8 50.3628 13177 13156 50.4 40.9 1.5 139 73.9 46 31 54.1 107 1127 19800 19917 50.4 50.9 50.2 40.9 0.2 117 71.7 42.7 31 50.3 109 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50.0 0.5 662 75.6 40.8 31 53.1 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 43.2 31 54.1 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 43.2 31 54.1 111 1132 27365 27385 53.2 47.8 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 113 1133 24448 24438 50 47.4 3633 24527 24508 50.5 45.0 50.5 45.0 51.0 71.3 42.7 31 50.3 113 13 24418 24438 50 47.4 3633 24527 24508 50.5 44.0 0.5 758 75.8 41.1 31 53.7 115 135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 33.7 3415 33.7 33.7 33.7 33.7 33.7 33.7 33.7	99	1119	8221	8240	52.4	50 3619	8929	8911	53.4	52.6	1	.709	75.4	40.1	30	53.6
102 1122 19709 19730 51.3 40.9 3622 19921 19900 51.8 45.5 0.5 213 73.9 41.8 31 52.2 103 1123 16366 16386 54.4 52.4 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.2 104 1124 3 21 53.4 52.6 3624 256 235 52.6 45.5 0.8 24.5 76.1 46.1 31 54.2 105 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13058 51.8 50 3626 13177 13166 50.4 40.9 1.5 139 73.9 46 31 52.2 107 1127 19800 19817 50.4 50.9 5627 19916 19895 50.2 40.9 0.2 117 71.7 42.7 31 50.3 108 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50 0.5 662 75.6 40.8 31 53.1 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 111 131 339 13058 51.8 50.8 50.8 50 50.8 40.9 0.4 709 76.6 43.2 31 54.1 111 131 3 27355 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.8 13 131 32 24418 24436 50 47.4 3633 24527 24508 50.5 44.4 0 756 75.9 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8	100	1120	13039	13057	51.1	52.6 3620	13177	13156	50.4	40.9	0.7	139	73.9	46	31	52
103 1123 16366 16386 54.4 52.4 3623 16774 16751 53.6 41.7 0.8 409 75.1 41.1 31 53.8 104 1124 3 21 55.4 52.6 36524 256 235 52.6 45.5 0.8 254 76.1 46.1 31 53.8 106 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52 107 1127 19800 19817 50.4 50 3626 13177 13156 50.2 40.9 0.2 117 17.7 42.7 31 50.3 109 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50.2 40.9 0.2 117 71.7 42.7 31 50.3 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 111 131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54 112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 107 70.8 43 31 50.6 131 133 24418 24430 50 47.4 3633 24527 24508 50.5 40.5 51 107 71.3 42.7 31 50.8 114 1134 26708 26727 50 45.3 340.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 155 155 141 135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41.1 31 53.7 155	101	1121	19801	19819	53.2	52.6 362	19917	19895	52.5	43.5	0.8	'117	72	-43.6	31	51.2
104 1124 3 21 53.4 52.6 3624 256 235 52.6 45.5 0.8 254 76.1 46.1 31 54.2 105 1125 4 22 52.3 52.6 3625 314 296 50.6 47.4 1.7 311 76.8 46.6 31 54.1 106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52 107 1127 19800 19817 50.4 50 3627 19916 19895 50.2 40.9 0.2 117 71.7 42.7 31 50.3 108 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50 0.5 662 75.6 40.8 31 53.1 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.2 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.2 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54.2 112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 13 13 3 24448 24436 50 47.4 3633 24527 24508 50.5 45 50.5 110 71.3 42.7 31 50.8 114 1134 26708 26727 50.4 50.8 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	102	1122	19709	19730	51.3	40.9 3622	19921	19900	51.8	45.5	0.5	213	73.9	41.8	31	
105 1125	103	1123	16366	16386	54.4	52.4 3623	16774	16751	53.6	41.7	0.8	.409		41.1	31	53.8
106 1126 13039 13058 51.8 50 3626 13177 13156 50.4 40.9 1.5 139 73.9 46 31 52 107 1127 19800 19817 50.4 50 3627 19916 19895 50.2 40.9 0.2 117 71.7 42.7 31 51.3 109 1128 4645 4655 50.2 42.9 3628 53.06 50.2 40.9 0.2 117 71.7 42.7 31 51.3 1109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 110 113 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54 111 1131 3 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 131 133 24418 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 50 114 1134 26708 26727 50 45 3634 27463 27463 27465 50 44.4 0.7 56 75.9 41.1 31 53.7 53.2 4179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 411 31 53.7	104	1124	3	21	53.4	52.6 3624	256	235	52.6	45.5	0.8	254	76.1	46.1	31	54.2
107 1127 19800 19817 50.4 50 3627 19916 19895 50.2 40.9 0.2 117 71.7 42.7 31 50.3 109 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50 0.5 662 75.8 40.8 31 53.1 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 1.1 709 76.6 43.2 31 54 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54 112 1132 27365 27365 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 51.1 1131 32 24148 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 53.7 50.8 40.9 41.1 40.7 56 75.9 41.1 31 53.2 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7 53.7 53.7 53.7 53.8 53.8 53.8 53.8 53.7 53.8 53.	105	1125	4	22	52.3	52.6 362	314	296	50.6	47.4	1.7	311	76.8	46.6	31	54.1
108 1128 4645 4665 50.2 42.9 3628 5306 5289 50.8 50 0.5 662 75.6 40.8 31 53.1 109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54.1 110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 1.1 709 76.6 43.2 31 54.1 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 43.2 31 54.1 112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 13 1133 24418 24439 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 50.8 114 1134 26708 26727 50 45 3634 27463 27446 50 44.4 0 768 75.8 75.8 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	106	1126	13039	13058	51.8	50 3626	13177	13156	50.4	40.9	1.5	139	73.9	46	31	52
109 1129 13039 13057 51.1 52.6 3629 13747 13726 50.8 40.9 0.4 709 76.6 43.2 31 54 100 130 3039 13058 51.8 50 3630 13747 13726 50.8 40.9 1.1 709 76.6 43.2 31 54 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54 112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 131 133 24418 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 50 50 114 1134 26708 26727 50 45 3634 27463 27464 50 44.4 0 756 75.9 41.1 31 53.2 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	107	1127	19800	19817	50.4	50 362	19916	19895	50.2	40.9	0.2	117	71.7	42.7	31	50.3
110 1130 13039 13058 51.8 50 3630 13747 13726 50.8 40.9 1.1 709 76.6 43.2 31 54 111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54 112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 113 1133 24418 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 70.3 42.7 31 50 114 1134 26708 26727 50 45 3634 27463 27446 50 44.4 0 756 75.9 41.1 31 53.7 115 1135 24179 24200 53.3	108	1128	4645	4665	50.2	42.9 362	5306	5289	50.8	50	0.5	662		40.8	31	53.1
111 1131 3 21 53.4 52.6 3631 253 233 51.8 47.6 1.6 251 76.2 46.2 31 54 112 1132 27365 27365 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 113 1133 24418 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 50 114 1134 26708 26727 50 45 3634 27463 27446 50 44.4 0 756 75.9 41.1 31 53.7 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	109	1129	13039	13057	51.1	52.6 3629	13747	13726	50.8	40.9	0.4	709	76.6	43.2	31	54
112 1132 27365 27385 53.2 47.6 3632 27464 27444 53 42.9 0.2 100 70.8 43 31 50.6 113 1133 24418 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 50.6 114 1134 26708 26727 50 445 3634 27463 27464 50 44.4 0 75.9 41.1 31 53.2 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 75.8 75.8 41.1 31 53.7	110	1130	13039	13058	51.8	50 3630	13747	13726	50.8	40.9	1.1	709	76.6	43.2	31	54
113 1133 24418 24436 50 47.4 3633 24527 24508 50.5 45 0.5 110 71.3 42.7 31 50 114 1134 26708 26727 50 45 3634 27463 27446 50 44.4 0 756 75.9 41.1 31 53.2 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	111	1131	3	21	53.4	52.6 363	253	233	51.8	47.6	1.6	251	76.2	46.2	31	54
114 1134 26708 26727 50 45 3634 27463 27446 50 44.4 0 756 75.9 41.1 31 53.2 115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	112	1132	27365	27385	53.2	47.6 363	27464	27444	53	42.9	0.2	100	70.8	43	31	50.6
115 1135 24179 24200 53.3 40.9 3635 24936 24919 51.8 50 1.5 758 75.8 41 31 53.7	113	1133	24418	24436	50	47.4 363	24527	24508	50.5	45	0.5	110	71.3	42.7	31	50
	114	1134	26708	26727	50	45 363	27463	27446	50	44.4	0	756	75.9	41.1	31	53.2
116 1136 26708 26727 50 45 3636 27462 27444 50.1 42.1 0.1 755 75.9 41.2 31 53.2	115	1135	24179	24200	53.3	40.9 363	24936	24919	51.8	50	1.5	758	75.8	41	31	53.7
	116	1136	26708	26727	50	45 363	27462	27444	50.1	42.1	0.1	755	75.9	41.2	31	53.2

	7 1137	26708		-		3637	27465				0.4	758	75.9	41.	3 3	1 54.5
111		27365					27464	27446	51.7	47.4	0.9	100				
119		27365				3639	27464	2744	52.4	45	0.2	100	70.8			
120		27365				3640	27464	2744	53	42.9	0.4	100				
12		27367		51.4	52.6	3641	27571	27552	50.1	40	1.3	205				
122		27367	-	51.4			27567	27547	51.1	42.9	0.2	201	74.7	44.		
123	1	2427					3186	3165	50.4	40.9	1.7	760		4	_	53.9
124		8867	8887	52.3		3644	9256	9237	50.8	45	1.5	390	75.1	41.		52.9
125		9934				3645	10605	10588	51.1	50	0.4	672				53.4
126		2429		50.2	47.4	3646	3186	3165	50.4	40.9	0.2	758			_	53.8
127		27365	27385	53.2	47.6	3647	27464	27445	52.4	45	0.8	100				50.4
128	-	19994	20011	50.4	50	3648	20615	20597	50.6	47.4	0.2	622	75.2	40	_	52.9
129	1	9922	9941	51.3	. 50	3649	10605	10588	51,1	50	0.2	684	75.8	41.2	-	53.5
130		12962	12980	50.7	47.4	3650	13544	13525	52.6	55	1.8	583	77.5	45.6	_	54.5
131		12965	12988	54	41.7	3651	13544	13525	52.6	55	1.5	580	77.4	45,5		55
132		· 13176	13197	52.7	45.5	3652	13544	13525	52.6	55	0.1	369	77.1	46.3		54.8
133		28867	28886	53.2	50	3653	29298	29280	51.4	52.6	1.7	432	76.8	45.1	32	54.8
134	1154	24418	24439	52.9	45.5	3654	25182	25164	51.4	47.4	1.5	765	76.1	41.7	32	53.8
135		24420	24440	50.8	42.9	3655	25182	· 25164	51.4	47.4	0.6	763	76.1	41.7		53.6
136	1156	8867	8887	52.3	47.6	3656	9107	9086	51.6	45.5	0.7	241	74.1	41.5		52.5
137	1157	1402	1422	50.2	42.9	3657	2103	2083	50.6	42.9	0.4	702	76.7	43.3		
138	1158	25782	25805	52.1	41.7	3658	26183	26163	51.7	42.9	0.4	402	74.7	40.3	32	53.8
139	1159	25781	25805	53.5	40	3659	26183	26160	54.5	41.7	1	403	74.7	40.3	32	52.9 53.4
140	1160	25781	25805	53.5	- 40	3660	26183	26159	54.9	40	1.5	403	74.7	40.2	32	53.4
-141	1161	2671	2692	52.1	40.9	3661	. 3052	3033	50.3	50	1.8	382	74.8	40.6		52.5
142	1162	12726	12746	51.3	47.6	3662	13177	13156	50.4	40.9	0.9	452	76.4	43.8	32	-53.7
143	1163	16909	16928	50.8	45	3663	17111	17090	51.1	40.9	0.3	203	75	44.8	32	52.8
144	1164	12234	12252	50.6	47.4	3664	12993	12975	51.4	47.4	0.8	760	76.4	42.5	32	53.8
145	1165	26039	26057	52.6	52.6	3665	26828	26810	52.9	52.6	0.2	790	76.4	42.4	32	54.4
	1166	26039	26057	52.6	52.6	3666	26694	26677	51.4	50	1.2	656	75.7	41	32	53.5
147	1167	26039	26057	52.6	52.6	3667	26692	26674	51.9	. 52.6	0.7	654	75.7	41	32	53.6
	1168	26039	26057	52.6	52.6	3668	26691	26673	51.3	47.4	1.3	653	75.6	40.9	32	53.4
149	1169	26039	26057	52.6	52.6	3669	26687	26669	51.3	47.4	1.3	649	75.6	40.8	32	53.4
150	1170	26039	26057	52.6	52.6	3670	26684	26666	53.4	52.6	0.8	646	75.6	40.9	32	53.8
151	1171	26039	26057	52.6	52.6	3671	26683	26665	52.7	52.6	0.1	645	75.6	40.9	32	53.8
	1172	9934	9953	50.7	50	3672	10449	10431	50.9	47.4	0.2	516	75.4	40.9	32	53.1
	1173	9927	9945	50.8		3673	10455	10434	51.1	40.9	0.3	529	75.3	40.9	32	53.1
	1174	7728	7746	51.7	52.6	3674	8188	8169	50.5	45	1.2	461	75.6	41.9	32	53.2
	1175	18550	18571	50.4	40.9	3675	19216	19195	50.2	40.9	0.2	667	75.7	41.1	32	53.2
	1176	19801	19819	53.2	52.6	3676	19921	19899	52.4	43.5	0.8	121	72.3	43.8	32	51.4
	1177	19709	19730	51.3	40.9	3677	19923	19904	50.1	50	1.2	215	73.9	41.9	32	51.4
	1178	4639	4659	51.1	47.6	3678	5306	5289	50.8	50	0.3	668	75.6	40.9	32	53.3
	1179	19794	19813	50	50	3679	19921	19901	50.2	47.6	0.2	128	72.6	43.8	32	50.9
	1180	12965	12985	51.2	42.9	3680	13544	13525	52.6	55	1.4	580	77.4	45.5	32	
	1181	9932	9953	53	45.5	3681	10449	10425	54.6	40	1.6	518	75.3	40.7	32	54.6
	1182	19795	19814	50.4	45	3682	19921	19900	51.8	45.5	1.4	127	72.3	43.3	32	
	1183	27366	27384	52.2		3683	27468	27451	51.1	50	1	103	71.3	43.3	32	50.9
	1184	27366	27384	52.2	52.6	3684	27467	27450	52.1	50	0.1	102	71.4	44.1		
165	1185	27366	27384	52.2	52.6		27466	27449	51	50	1.2	101	71.5	44.1	32	50.7
166	1186	25782	25805	52.1		3686	26183	26164	51	45	1.1	402	74.7	44.6	32	50.4
	1187	9934	9953	50.7	50		10449	10428	51.9	40.9	1.2	516	75.4	40.3	32	52.7
168	1188	9925	9945	53.4		3688	10449	10425	54.6	40.9	1.2	525	75.4	40.9	32	53.1
								+20	54.0	+0	1.2	020	10.4	41	32	53.9

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	1189	19800	19817	50.4		3689	19922	19902	50	42.9	0.4	123	72.1	43.1	32	50.6
	1190	8867	8887	52.3	47.6		9310	9291	51.2	45	1.2	444	75.4	41.4	32	53.2
	1191	27367	27385	51.4	52.6		27468	27451	51.1	50	0.3	102	71.4	44.1	32	50.4
172	1192	27367	27385	51.4	52.6	3692	27467	27450	52.1	50	0.7	101	71.5	44.6	32	50.6
173	1193	2671	2692	52.1	40.9	3693	3082	3058	52.3	40	0.2	412	74.9	40.5	32	53.2
174	1194	9927	9945	50.8	52.6	3694	10608	10589	51	50	0.2	682	75.8	41.2	32	53.4
175	1195	19800	19817	50.4	50	3695	19920	19899	50.2	40.9	0.3	121	71.9	43	32	50.5
176	1196	13177	13197	50.3	42.9	3696	13547	13528	50.2	45	0.1	371	76.9	45.8	32	54
177	1197	28179	28200	50.8	40.9	3697	28672	28654	50.6	52.6	0.3	494	79.8	51.8	32	56.1
178	1198	27367	27385	51.4	52.6	3698	27466	27449	51	50	0.4	. 100	71.6	45	32	50.5
179	1199	27366	27385	52.8	50	3699	27465	27446	54.6	50	1.7	100	71.2	44	32	50.8
180	1200 ·	19800	19818	52.1	52.6	3700	19921	19901	50.2	47.6	2	122	72.2	43.4	32	50.7
181	1201	9927	9945	50.8	52.6	3701	10455	10435	50.5	42.9	0.3	529	75.3	40.6	32	52.9
182	1202	-28868	28887	50.7		3702	29298	29280	51.4	52.6	0.7	431	76.8	45	32	54.1
183	1203	28867	28887	53.7	47.6	3703	29306	29288	53.5	52.6	0.3	440	76.9	45.2	32	55
184	1204	28867	28887	53.7	47.6	3704	29301	29282	55.3	55	1.5	435	76.9	45.3	32	55.1
185	1205	28868	28888	51.4	42.9	3705	29298	29280	51.4	52.6	0	431	76.8	45	32	54.3
186	1206	28867	28888	54.3		3706	29306	29288	53.5	52.6	0.8	440	76.9	45.2	32	55
187	1207	28867	28888	54.3	45.5		29301	29282	55.3	55	1	435	76.9	45.3	32	55.2
188	1208	28870	28889	50.1		3708.	29298	29280	51.4	52.6	1.3	429	76.8	45	32	53.9
189	1209	28868	28889	52	40.9	3709	29306	29288	53.5	52.6	1.5	439	76.9	45.1	32	54.5
190	1210	28867	28889	54.8		3710	29301	29282	55.3	55	0.5	435	76.9	45.3	32	55.4
191	1211	28867	28890	55.2		3711	29306	29288	53.5	52.6	1.7	440	76.9	45.2	32	55
192	1212	28867	28890	55.2		3712	29301	29282	55.3	55	0.1	435	76.9	45.3	32	.55.5
193	1213	28867	28890	55.2	41.7		29299	29280	53.9	55	1.3	433	76.9	45.3	32	55.1
		12234	12252	50.6	47.4		12996	12977	50.2	40	0.3	763	76.4	42.3	32	53.6
195	1215	28968	28988	50.9	47.6	3715	29298	29280	51.4	52.6	0.6	331	76.2	44.7	32	53.7
196	1216	28968	28989	51.5		3716	29298	29280	51.4	52.6	0.1	331	76.2	44.7	32	53.9
197	1217	13230	13251	52.4		3717	13544	13525	52.6	55	0.1	315	77.2	47.3	32	54.8
198	1218	29186	29205	50.1		3718	29298	29280	51.4	52.6	1.3	113	72.8	46	32	51.1
199	1219	29195	29213	51.9	52.6	3719	29306	29288	53.5	52.6	1.6	112	73.6	48.2	32	52.2
200	1220	29195	29213	51.9	52.6	3720	29298	29280	51.4	52.6	0.5	104	73.1	48.1	32	51.7
201	1221	29196	29214	51.1	52.6		29298	29280	51.4	52.6	0.3	103	73.3	48.5	32	51.7
202	1222	29195	29214	52.6	50	3722	29306	29288	53.5	52.6	0.9	112	73.6	48.2	32	52.4
203	1223	29196	29215	51.8	50	3723	29306	29288	53.5	52.6	1.6	111	73.8	48.6	32	52.3
204	1224	29196	29215	51.8	50	3724	29298	29280	51.4	52.6	0.4	103	73.3	48.5	32	51.8
205	1225	29197	29216	50	45	3725	29298	29280	51.4	52.6	1.4	102	73	48	32	51.2
206	1226	29196	29216	52.5	47.6	3726	29306	29288	53.5	52.6	1	111	73.8	48.6	32	52.5
207	1227	29195	29216	53.8	45.5	3727	29301	29282	55.3	55	1.5	107	73.5	48.6	32	52.7
208	1228	29254	29273	53.1	50	3728	29358	29339	52.8	50	0.2	105	73.4	48.6	32	52.3
209	1229	29259	29278	52.6	50	3729	29358	29339	52.8	50	0.2	100	72.4	47	32	51.6
210	1230	1402	1422	50.2	42.9	3730	1773	1755	51.7	52.6	1.5	372	75.8	43.3	33	53.2
211	1231	12726	12746	51.3	47.6	3731	13326	13306	50.7	42.9	0.6	601	76.7	43.6	33	. 54
212	1232	4	22	52.3	52.6	3732	269	251	51.1	52.6	1.2	266	76.5	46.6	33	54
213	1233	19800	19817	50.4		3733	19923	19903	50.9	47.6	0.4	124	72.3	43.5	33	50.9
214	1234	2371	2389	50.3	47.4	3734	3082	3058	52.3	40	2	712	76.7	43.3	33	53.9
215	1235	3	21	53.4		3735	270	251	52.9	50	0.5	268	76.4	46.3	33	54.4
216		9930	9949	52.2		3736	10183	10166	50.9	50	1.3	254	75.3	44.1	33	53.1
217	1237	19795	19814	50.4		3737	19923	19904	50.1	50	0.3	129	72.5	43.4	33	50.9
218	1238	8867	8887	52.3	47.6		9365	9347	53	52.6	0.7	499	75.8	42.1	33	53.9
219		2371	2389	50.3	47.4		3055	3036	50.6	50	0.3	685	76.7	43,4	33	53.9
						_,			- 00.0		U.U	555		.0.7	_~	

r																	
Ţ		0 1240	19709			3 40.9	3740	19921	1990	50.2	47.6	1.1	213	73.9	41.8	33	51.9
- 1	22	1 1241	9930	9949	52.2	2 50	3741	10183	1016				_				
ſ	22	2 1242	2371	2389	50.3	47.4	13742	2747									
- [22	3 1243	24921	24938	50.4		3743	25182				1		76.9		1	54
ı	22	4 1244	18077	18099			3744	18443					262	74.2	41.2	_	
ı	22		25772					26183				1.5		75.8	43.3		54.5
ı	22		25769			_	3746	26183			45	1.3		74.8	40.3		52.8
ŀ	22		25348				3747					0.8		74.9	40.5		52.6
ŀ	22		12726					25548			50	0.1	201	74.3	43.3	33	52.4
ŀ	22		8372				3748	13323			45	0.2	598	76.7	43.6	33	54.1
ŀ	23			8390			3749	8928		51.9		1.2	557	75.1	40	_33	52.9
H	23		2671	2692			3750	3189			45.5	1.2	519	75.7	41.6	33	53.4
ŀ			25348	25365			3751	25548		51.1	50	0.7	201	74.3	43.3	33	52.2
ŀ	23		19801	19819	53.2		3752	19923	19902	51.5	45.5	1.7	123	72.4	43.9	33	51.3
H	233		27442	27461	51.5		3753	27546	27527	51.3	50	0.2	105	71.8	44.8	33	50.8
ŀ	234		8867	8887	52.3		3754	9312	9293	50.6	45	1.8	446	75.4	41.5	33	53
1	235		2671	2692	52.1		3755	3056	3038	50.8	52.6	1.3	386	74.8	40.7	33	52.7
L	236		13231	13251	50.1		3756	13547	13528	50.2	45	0.2	317	76.9	46.7	33	54
L	237		9055	9079	52.8	40		9310	9291	51.2	45	1.7	256	74.4	41.8	33	52.5
L	238		28821	28838	50.3	50	3758	29298	29280	51.4	52.6	1.1	478	77	45.2	33	54.1
L	239		9055	9079	52.8	40	3759	9253	9235	51.6	47.4	1.2	199	73.6	41.7	33	52.1
L	240	1260	23840	23863	55.2	45.8	3760	24050	24031	56.5	55	1.4	211	75.0	44.5	33	54.1
	241	1261	18074	18093	50.3	45	3761	18233	18214	52	50	1.7	160	73.9	44.5	33	
	242	1262	27366	27384	52.2	52.6	3762	27674	27654	51.9	42.9	0.3	309	74.3			. 51.9
Γ	243	1263	28967	28989	53.7	47.8	3763	.29301	29282	55.3	55	1.5	335	76.4	40.5	33	52.7
Γ	244	1264	27366	27384	52.2		3764	27674	27653	52.5	40.9	0.3	309	74.3	45.1	33	54.7
E	245	1265	28966	28988	55.3		3765	29301	29282	55.3	55	0.1	336	76.4	40.5	33	52.8
Г	246	1266	18074	18094	51.1	42.9	3766	18233	18214	52	50	1	160	73.9	45.2 44.4	33	55.2
	247	1267	28965	28984	52.9		3767	29298	29280	51.4	52.6	1.5	334	76.4	45.2	33	52.1
Г	248	1268	18081	18099	51.2	52.6	3768	18233	18215	51.3	52.6	0.1	153	74	45.1	33	52.2
L	249	1269	18081	18099	51.2	52.6	3769	18233	18214	52	50	0.8	153	74	45.1	33	52.2
L		1270	18081	18099	51.2	52.6	3770	18231	18210	52.2	45.5	1	151	73.6	44.4	33	
	251	1271	24480	24500	53.2	47.6	3771	24815	24791	54.5	40	1.3	336	75.6			. 52
Г	252	1272	24481	24503	52.7	43.5	3772	24815	24791	54.5	40	1.8	335	75.5	43.2	33	54
Г	253	1273	27367	27385	51.4	52.6		27675	27656	50	40	1.4			43	33	53.8
Г	254	1274	27367	27385	51.4	52.6		27674	27654	51.9	42.9	0.5	309	74.3	40.5	33	52.1
Γ	255	1275	27367	27385	51.4	52.6		27674	27653	52.5	40.9		308	74.4	40.6	33	52.6
	256	1276	18081	18099	51.2	52.6		18223	18206	51.8	50	1.1	308	74.4	40.6	33	52.6
r	257	1277	18080	18099	53		3777	18220	18202	51.8		0.6	143	73.2	44.1	33	51.7
Г	258	1278	9933	9952	50.9		3778	10670	10649	51.3	52.6 40.9	1.9	141	73.1	44	33	52.2
r	259	1279	27665	27686	51.4		3779	28208	28190			0.5	738	75.7	40.8	33	53.4
Γ	260	1280	27665	27685	50.7		3780	28208	28190	51.7 51.7	52.6 52.6	0.4	544	75.1	40.1	33	53.1
Г		1281	27442	27461	51.5		3781	27541	27522	50.1	45	1.1	544	75.1	40.1	33	52.9
Г	262	1282	28821	28840	51.8		3782	29298	29280			1.4	100	71.2	44	33	50
Г	263	1283	28821	28839	51.1	47.4		29298	29280	51.4	52.6	0.4	478	77	45.2	33	54.4
r	264	1284	8868	8889	50.4		3784	9252			52.6	0.3	478	77	45.2	33	54.3
Г		1285	19800	19818	52.1		3785	19920	9235 19899	50.1	50	0.3	385	75.1	41.3	34	52.7
r		1286	9055	9079	52.8		3786	9313			40.9	2	121	71.9	43	34	50.5
H		1287	10142	10163	51.3		3787	10605	9293 10588	52.1	47.6	0.7	259	74.6	42.1	34	52.9
Н		1288	12726	12746	51.3	47.6		13312		51.1	50	0.2	464	74.9	40.1	34	52.8
Г		1289	9055	9079	52.8	40 3		9257	13294	51	52.6	0.3	587	76.6	43.6	34	54
Г		1290	7876	7895	51.5	45 3		8188	9237	52.2	42.9	0.7	203	73.5	41.4	34	52.2
				. 555	01.0	+510	11 30	0188	8169	50.5	45	1.1	313	75	42.2	34	52.8

271	1291	23843	23863	50.3	420	3791	24527	24507	51	42.9	0.7	1 005	70			
272	_	10247	10267	50.5		3792	10608	10589	51	50		685	76	41.8	34	53.4
273		24179	24199	52.7			24815	24791	54.5	40	0.5	362	74.6	40.3	34	52.4
274		12236	12256	51.2							1.8		75.8	41.3	34	53.9
275		7869	7889	52.5		3795	12998	12979	50.1	45	1.1	763	76.4	42.5	34	53.6
276					_		8189	8169	52	47.6	0.5		75.3	42.7	34	53.4
277		1402	1422	50.2	_		2152	2133	50.7	45	0.5	751	76.7	43.1	34	53.8
	1297	12233	12251	51.1		3797	12993	12975	51.4	47.4	0.2	761	76.5	42.6	34	54
278		3033	3053	51.7		3798	3650	3631	53.1	50	1.4	618	76.4	42.9	34	54.1
279		12233	12251	51.1		3799	12996	12977	50.2	40	0.9	764	76.4	42.4	34	53.7
280	1300	24483	24503	51	42.9		24938	24921	50.4	50	0.6	456	75.6	41.9	34	53.1
281	1301	11541	11561	50.9	42.9	3801	12253	12235	50.1	52.6	0.8	713	76.2	42.1	34	53.5
282		24622	24643	57.1	54.5	3802	25400	25379	56	50	1.1	779	75.7	40.7	34	54.9
283	1303	24622	24643	57.1	54.5	3803	25400	25378	56.4	47.8	0.6	779	75.7	40.7	34	55
284	1304	24630	24648	50.8	52.6	3804	25403	25385	51.1	47.4	0.3	.774	75.7	40.6	34	53.3
285	1305	9929	9946	50	50	3805	10605	10588	51.1	50	1	677	75.8	41.2	34	53.2
286	1306	24633	24651	50.1	52.6	3806	25403	25385	51.1	47.4	1	771	75.6	40.5	34	53.1
287	1307	11541	11560	50.1	45	3807	12253	12235	50.1	52.6	0	713	76.2	42.1	34	53.5
288	1308	24635	24653	50.5	52.6	3808	25403	25385	51.1	47.4	0.7	769	75.6	40.4	34	53.2
289	1309	9933	9952	50.9	45	3809	10608	10589	51	50	0.1	676	75.8	41.1	34	53.4
290	1310	24921	24938	50.4	50	3810	25548	25531	51.1	50	0.7	628	75.6	40.9	34	53.1
291	1311	7725	7743	50.8	47.4		8188	8169	50.5	45	0.4	464	75.6	41.8	34	53.2
292	1312	28547	28568	53.5	45.5	3812	29301	29282	55.3	55	1.8	755	78.5	47.5	34	56.1
293	1313	28547	28568	53.5	45.5		29306	29288	53.5	52.6	0	760	78.5	47.5	34	56.1
294	1314	28548	28568	50.5	42.9	3814	29298	29280	51.4	52.6	0.9	751	78.4	47.4	34	55.2
295	1315	28546	28567	55.1		3815	29301	29282	55.3	55	0.2	756	78.5	47.6	34	56.6
296	1316	28547	28567	52.9	47.6		29298	29280	51.4	52.6	1.5	752	78.5	47.5	34	55.5
297	1317	28546	28565	52.2	50		29298	29280	51.4	52.6	0.8	753	78.5	47.5	34	55.5
298	1318	28546	28565	52.2		3818	29306	29288	53.5	52.6	1.3	761	78.5	47.6	34	55.7
299	1319	28396	28416	52.4		3819	28672	28654	50.6	52.6	1.8	277	78.6	51.6	34	
300	1320	28396	28415	51.2	45	3820	28671	28652	52.8	55	1.6	276	78.6			55.3
301	1321	28396	28415	51.2	45		28671	28653	50.2	52.6	1.0	276	78.6	51.4	34	55.4
302	1322	12976	12995	51.1	45		13545	13527	50.2	52.6	0.8	570	77,4	51.4 45.6	34	55.2
	1323	16551	16568	51.1	50		16711	16691	51	42.9	0.8	161	73.8	44.1		54.4
304	1324	28395	28414	51.5		3824	28672	28654	50.6	52.6	0.9	278	78.6	51.4	34	55.3
_	1325	16555	16572	50.3	50	3825	16780	16760	51.4	42.9	1.1	226				
	1326	28394	28413	51.8	45	3826	28671	28652	52.8	55	1.1	278	73.6	40.7	34	51.7
307	1327	28395	28413	50.2	42.1	3827	28671	28653	50.2	52.6			78.6	51.4	34	55.6
	1328	7728	7746	51.7	52.6	3828	8049	8032	50.4		0	277	78.5	51.3	34	55.1
	1329	28394	28412	51.1	47.4	3829	28671	28652	52.8	50 55	1.3	322	74.9	41.6	34	52.6
	1330	28394	28412	51.1	47.4	3830	28671	28653	50.2	52.6	0.8	278	78.6	51.4	34	55.4
311	1331	11543	11562	50.4	40	3831	12257	12237	51.3	47.6	0.8	278	78.6	51.4	34	55.2
	1332	28393	28411	52.9	52.6	3832	28671	28652	52.8			715	76.2	42	34	53.5
	1333	28394	28411	50.3	50	3833	28671	28653	50.2	55	0.1	279	78.6	51.6	34	56
	1334	4255	4276	51.7	45.5	3834	4710			52.6	0	278	78.6	51.4	34	55.2
	1335	12975	12994	52.1		3834	13545	4691	50.2	45	1.5	456	75.1	40.8	34	52.8
	1336	9930	9948	51.5				13526	52.9	55	0.8	571	77.4	45.5	34	54.9
	1337	27665	27686			3836	10608	10589	51	50	0.5	679	75.8	41.2	34	53.5
	1338	27665	27686	51.4		3837	28411	28393	52.9	52.6	1.6	747	76.8	43.5	34	54.3
	1338	11541	11561	51.4		3838	28415	28396	51.2	45	0.2	751	76.8	43.4	34	54.2
	1340	27665		50.9		3839	12257	12237	51.3	47.6	0.5	717	76.2	42	34	53.7
	1340	11543	27685 11562	50.7		3840	28415	28396	51.2	45	0.5	751	76.8	43.4	34	54.1
321	1041	11043	11002	50.4	40]	3841	12253	12235	50.1	52.6	0.3	711	76.2	42.1	34	53.5

														1,002		1,10	
3	22 13	12 11	545 1	1563	3 50.	8 47	.4 384	2 122	54 122	26 50	0.5 47	41 0	al =:				
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33				2243								5 0.4		4 75.6	41.	B 34	53
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33				9953	50.7		_					0 1.2	, ,,,		41.2	2 34	53.5
33				2243	50.2	_	0 3853					9 0.3	52	75.3	40.6	3 34	
33				3815		-		267			.9 52.	6 0.7	450	77	45.3	3 34	
33					50.9		-	444			.6 42.	9 0.4	649	75.4	40.4		
33				953	50.7		-1	1045		-1	1 40.	9 0.4	522	75.3	40.6		
33				3093	50.3			1869			9 52.	6 1.5	624	76.2	42.5		
33				2994	50.3			1354		7 50.	3 52.	6 0.1	570		45.6		54.4
34				2057	50.6	_	-10000	1249		0 5	0 47.	4 0.6	459		43.6		53.5
34				2057	50.6	_	1-0-0	1225			3 47.	0.7	218	75.4	45.4	_	53.1
34				557	50.4			1225		7 51.	3 47.0	0.9	718	76.2	42.1	34	53.6
34				993	51.4			1354		52.	9 5	1.5	571	77.4	45.5	34	54.7
34				993	51.4			1354			3 52.6	1.1	571	77.4	45.5	34	54.4
34	-1			560	53.2	47.6		11983			3 52.6	0.2	444	75.1	40.8	34	53.6
34				057	50.6	50	-	12253			52.6	0.5	214	75.5	45.8	34	53
34		1303		995 057	51.1	45	1	13545			9 55	1.8	570	77.4	45.6	34	54.6
34				380	51.1	52.6	1000	13314			50	0.1	276	75.7	44.6	34	53.4
349				380	52.4	55		27463			40	0.8	103	71.7	44.7	34	50.7
350				380	52.4		3869	27463		- 00.0		1.6	103	71.7	44.7	34	50.5
35	1.0.0	2534		365	52.4 50.4	55		27464				0.7	104	71.9	45.2	34	51
352				941	51.3		3871	25645		50.8		0.4	298	74.6	41.3	34	52,4
	3 1373	1303			51.3	50		10449		50.9		0.3	528	75.4	40:9	34	53.2
354		1223			50.1		3873	13323	13304	51.1		0	285	75.8	44:6	34	53.5
355		301	_	36	50.1		3874	12412	12392	50		0.1	178	73.2	41.6	34	51.3
356		1303			51.1		3875	3185	3164	51	45.5	0.7	170	74.5	45.3	34	52.3
357		786		89	52.5			13326	13306	50.7	42.9	0.4	288	75.8	44.4	34	53.4
358		2642			51.5		3877 3878	8050	8032	52	52.6	0.5	182	73.8	42.9	34	52.3
359		2642			51.5		3879	26655	26634	50.6	40.9	0.9	235	74.1	41.7	34	52.2
360		2604			56.4	54.5	3880	26657	26639	50.8	47.4	0.7	237	74.2	41.8	34	52.3
361	1381	2604			56.4		3881	26183	26159	54.9	40	1.5	144	72	41	34	52
362	1382	2604			56.4		3882	26183	26160	54.5	41.7	2	144	72	41	34	51.9
363	1383	1237			50.8		3883	26184 12724	26161	55.1	41.7	1.3	145	71.9	40.7	34	52
364	1384	2604			56.4		3884		12705	52.4	55	1.6	352	75.6	42.9	34	53.2
365	1385	26039			54		3885	26589 26183	26569	54.7	47.6	1.7	550	75.1	40	34	54.1
366	1386	26039		_	54		3886		26159	54.9	40	0.9	145	71.9	40.7	34	51.7
367	1387	26039			54		3887	26183	26160	54.5	41.7	0.4	145	71.9	40.7	34	51.7
368	1388	26039			54		3888	26183 26184	26161	54	43.5	0	145	71.9	40.7	34	51.7
369	1389	26039			52.6		3889	26174	26163	53	40.9	1	146	71.8		34	51.3
370	1390	10246			50.4		3890	10605	26153 10588	51	40.9	1.6	136	71.8		34	50.7
371	1391	3234	32		51.1	47.6		3497	3478	51.1	50	0.6	360	74.5		34	52.4
372	1392	26039	260		52.6		3892	26183	26162	51.3	50	0.2	264	74.3		34	52.4
		·				32.5		20103	20102	52.8	45.5	0.2	145	71.9	40.7	34	51.2

	0=	1.00-	1 115	777	7		-										
-		3 1393	11540				3893	12253		50.1	52.6	0.3	714	76.2	42.2	34	53.5
\vdash	37		3234					3500	3481	51.2	50	0.1	267	74.3	41.2		52.4
L	37		3794		_			4445	4424	51.3	40.9	1.6	652	75.5	40.5		53.3
L		1396	3794		52.9	52.0	3896	4446	4425	51.8	45.5		653	75.5	40.6		53.5
	37		3234	3254	51.1	47.0	3897	3646	3625			1	413	75.1	41.2		53
	37		3234	3254	51.1	47.6	3898	3647	3628			0.5		75.2	41.3	34	52.9
	37	1399	3226	3245	51.7	56	3899	3497	3478			0.4	272	74.6	41.9	34	52.9
Г	38	1400	3797	3815	50.9	47.4	3900	4444	4424			0.4	648	75.4	40.4	34	53.1
	38	1401	3226	3245	51.7	55		3500	3481	51.2		0.5	275	74.6	41.8	34	52.7
	38	1402	16366	16384	50.3			16780	16760	51.4		1.1	415	75.1			
	383	1403	25782	25806	53.5	40		26183	26161	54		0.5	402		41	_34	52.7
\vdash	384		16366	16385	52.9	55		16780	16760	51.4	43.5	1.4	402	74.7 75.1	40.3	34	53.5
	385	1405	16367	16386	51.4	50		16781	16761	51.3		0.1			41	34	53.1
\vdash	386		12236	12256	51.2	42.9		12992	12974	51.2	52.6		415	75.1	41	34	53
\vdash	387		16367	16386	51.4	50		16777	16758			0	757	76.5	42.5	34	54
-	388		16367	16386	51.4	50		16711		51.5	50	0.1	411	75	40.9	34	53
	389		3226	3245	51.7	55			16691	51	42.9	0.3	345	75.2	42	34	53
		1410	16548	16566				3503	3484	51.5	50	0.3	278	74.7	42.1	34	52.9
_		1411	16549	16567	54.9		3910	16782	16760	54.3	43.5	0.6	235	74	41.3	34	53.2
	392		25354		54.9		3911	16782	16760	54.3	43.5	0.6	234	74	41.5	34	53.2
				25372	50.9	52.6		25645	25626	50.8	45	0.2	292	74.4	41.1	34	52.4
		1413	16551	16568	51.1		3913	17038	17021	50.7	50	0.4	488	75.8	42.2	34	'53.4
		1414	25348	25366	51.2	47.4		25645	25626	50.8	45	0.4	298	74.6	41.3	34	52.5
		1415	16551	16568	51.1	50		16780	16760	51.4	42.9	0.3	230	73.9	41.3	34	·52.2
-		1416	7725	7743	50.8	47.4		8049	8032	50.4	50	0.5	325	74.9	41.5	34	52.6
_	397		29200	29224	54.2	40		29299	29280	53.9	55	0.3	100	72.8	48	34	52.3
		1418	29200	29224	54.2		3918	29301	29282	55.3	55	1.1	102	73	48	34	52.4
_		1419	29200	29223	53.7	41.7		29299	29280	53.9	55	0.2	100	72.8	1 48	34	52.2
		1420	29200	29223	53.7		3920	29301	29282	55.3	55	1.6	102	73	· 48	34	52.3
_	401		29199	29222	54.6	41.7	3921	29301	29282	55.3	55	0.7	103	72.9	47.6	34	52.5
		1422	29200	29222	53.2	43.5	3922	29299	29280	53.9	55	0.7	100	72.8	48	34	52
		1423	29199	29221	54.1	43.5	3923	29301	29282	55.3	55	1.2	103	72.9	47.6	34	52.3
_	404		29200	29221	52.6	45.5	3924	29299	29280	53.9	55	1.3	100	72.8	48	34	51.9
	405	1425	18074	18093	50.3	45	3925	18239	18220	50	45	0.3	166	73.9	44	34	51.8
	406	1426	18074	18093	50.3	45	3926	18238	18219	50.3	45	0	165	74	44.2	34	51.9
	407	1427	1402	1426	54.1	40	3927	1774	1755	53.1	50	1	373	75.8	43.2	34	54.1
	408	1428	18074	18094	51.1	42.9	3928	18697	18679	51.9	52.6	0.8	624	76.2	42.5	34	53.8
	409	1429	18074	18094	51.1	42.9	3929	18239	18220	50	45	1	166	73.9	44	34	51.8
1	410	1430	18074	18094	51.1	42.9	3930	18238	18219	50.3	45	0.8	165	74	44.2	34	51.8
1	411	1431	3226	3245	51.7		3931	3504	3485	50.4	45	1.3	279	74.7	44.2	34	52.5
	412	1432	18081	18099	51.2		3932	18662	18641	50.4	40.9	0.7	582	76.3			
	413	1433	7725	7742	50		3933	8049	8032	50.4	50	0.7	325		42.8	34	53.6
		1434	29182	29205	54.6		3934	29301	29282	55.3	55	0.3		74.9	41.5	34	52.5
	415	1435	4255	4276	51.7		3935	4711	4692	51.2			120	73.4	46.7	34	52.9
_		1436	29183	29204	50.4		3936	29298	29280		45	0.5	457	75.1	40.7	34	53
	117	1437	3225	3243	50.9		3937	3497		51.4	52.6	1.1	116	72.8	45.7	34	51.2
		1438	29181	29202	53.9				3478	51.3	50	0.4	273	74.7	42.1	34	52.7
	119	1439	29182	29202	51.2		3938	29301	29282	55.3	55	1.4	121	73.6	47.1	34	52.8
		1440					3939	29298	29280	51.4	52.6	0.2	117	73.1	46.2	34	51.6
		1440	29180	29199	50.1		3940	29298	29280	51.4	52.6	1.3	119	73.2	46.2	34	51.4
			28970	28993	53.3		3941	29301	29282	55.3	55	1.9	332	76.1	44.6	34	54.4
	_	1442	28971	28993	51.9	43.5		29298	29280	51.4	52.6	0.5	328	76.1	44.5	34	53.8
		1443	4255	4276	51.7		3943	4711	4693	50.4	47.4	1.3	457	75.1	40.7	34	52.8
4	24	1444	12976	12996	51.8	42.9	3944	13545	13526	52.9	5 5	1.1	570	77.4	45.6	34	54.8
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426 1446 3226 3242 50.0 50.0 50.0 50.0 50.0 50.0 50.0 50.		-14	1														
427 1447 3225 3243 50.0 52.6 3947 30.03 3484 51.5 50 0.6 279 74.8 423 34 52.7 428 1449 3225 3243 50.0 52.6 3948 3504 3485 50.4 445 0.5 280 74.8 42.1 34 52.2 429 1449 3225 3243 50.0 52.6 3948 3504 2928 53.3 55.0 1.6 363 76.6 44.8 34 52.7 430 1450 28941 28961 51.6 42.9 3950 29206 62.3 55.3 55 0.1 363 76.6 44.8 34 54.1 431 1451 8867 8866 60.7 50 3951 9252 9285 50.1 50 0.6 386 75.1 41.5 34 52.7 432 1452 28939 28960 54.7 50 3952 28301 28282 55.3 55 0.1 367 76.5 44.8 34 54.1 433 1453 28941 28960 50.2 47.6 3952 2830 28280 55.3 55 0.6 386 76.1 41.5 34 52.4 433 1453 28941 28960 50.9 43 3954 22280 28280 51.4 52.6 0.5 586 76.3 41.5 34 55.4 435 1455 3380 3380 51.4 42.9 3955 3494 3473 50.4 40.9 1 135 73.7 45.9 34 51.8 435 1455 3380 3380 51.4 42.9 3955 3494 3473 50.4 40.9 1 135 73.7 45.9 34 51.8 435 1457 12372 12391 50.8 47.4 3957 12394 12376 50.3 47.4 0.4 62 76.4 42.9 34 53.4 436 1458 19794 19813 50 50 3958 19921 19900 51.8 45.5 1.8 122 72.6 43.8 34 50.9 440 1460 12234 12252 50.6 47.4 3957 12394 12376 50.3 47.4 0.4 62 76.4 42.9 43.5 34 441 1461 12234 12252 50.6 47.4 3960 12992 12776 50.3 47.4 0.2 76.1 64.8 34 53.4 441 1461 12234 12252 50.6 47.4 3960 12992 12776 50.3 47.4 0.2 76.1 64.8 34 53.8 442 1462 3034 3033 50.3 50 3662 3647 3686 1060 1068 51.1 50 0.6 680 75.1 41.2 34.5 52.8 443 1463 18794 19813 50.5 42.9 36.9								29306					339	76.3	44.8	3 34	54
428 448 3.225 3.243 50.0 52.6 3.044 3.004 3.485 50.4 45 0.5 207 74.8 42.3 34 52.6 429 1449 28939 28961 51.6 42.9 3905 29306 29286 53.5 55.0 1.8 366 76.4 44.8 34 52.6 431 1451 8867 8866 50.7 50.3 50.3 50.2 2282 53.5 52.6 1.8 366 76.4 44.8 5.2 431 1451 8867 8866 50.7 50.3 50.3 50.2 2282 53.5 52.6 1.8 366 76.4 44.8 14.5 34 52.7 431 1451 8867 8866 50.7 50.3 50.3 50.2 20282 53.5 52.6 1.8 366 76.6 44.8 14.5 34 52.7 432 1452 28933 22890 54.7 50.3 50.2 22801 22822 55.3 55.0 6.6 363 76.6 44.8 14.5 34 54.1 34 34.5						1	1				2 50	0.3	276	74.7	42	2 34	52.6
4429 4449 28889 28961 55.2 47.8 3849 28901 28926 55.3 55.0 1.8 365 76.4 44.8 34 54.1 34 55.2 44.0 1450 28941 28961 51.8 42.9 3950 29208 29.5 55.0 5.8 56.0 1.8 366 76.4 44.8 34 54.1 341 1451 8867 8866 60.7 50.9 3951 28222 29.25 50.1 50.0 50.8 366 76.4 44.8 34 54.1 34 44.1 44.1 44.1 44.1 28941 28960 52.4 47.6 3953 28901 28202 55.3 55.0 6.8 386 76.4 44.8 34 54.1 44.3 44.1 44.1 44.1 28941 28960 52.4 47.6 3953 28901 28208 53.5 55.0 6.8 38.6 76.5 44.5 34.5 44.5													279	74.8	42.3	3 34	52.7
430 1450 28941 28961 51.6 42.9 3950 29286 53.5 52.6 1.8 366 76.4 44.8 34 54.1 431 1451 8687 8868 50.7 50 3951 9252 9235 50.1 50 0.6 368 76.5 45.2 34 53.1 433 1453 228940 22806 54.7 50 3951 9252 9235 50.1 50 0.6 368 76.5 45.2 34 53.1 433 1453 228940 22806 52.4 47.6 3953 22920 2286 53.5 52.6 1 367 76.5 45.2 34 53.1 433 1453 228940 22806 52.4 47.6 3953 22920 22806 53.5 52.6 1 367 76.5 45.2 34 53.1 435 1455 3380 3380 51.4 42.9 3955 3494 3473 50.4 40.9 1 135 73.6 34.7 34 51.7 436 1456 19709 19730 51.3 40.9 3956 19910 18985 50.2 40.9 1 228 73.6 41.3 34 51.7 437 1457 12373 12393 50.8 47.4 3957 12994 12976 50.3 47.4 0.4 622 76.4 42.9 34.5 34.4 439 1459 3361 3381 50.5 42.9 3959 3494 3473 50.4 40.9 0.1 135 73.6 41.3 34 51.7 431 1451 1224 12252 50.6 47.4 3957 12990 12976 50.3 47.4 0.4 62.7 76.4 42.8 34 53.7 442 1460 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.4 62.7 64.4 42.8 34 53.7 443 1464 12224 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.4 62.7 64.4 42.8 34 53.7 444 1464 12224 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 761 76.4 42.4 34 53.7 445 1468 12977 12986 50.2 40.9 3863 3254 9226 50.8 47.4 1.7 388 75.1 43.1 34 53.7 446 1468 12977 12986 50.2 40.9 3967 3646 3625 50.5 47.4 1.2 29 75.1 43.1 34 53.6 446 1468 12977 12986 50.2 40.9 3967 3646 3625 50.5 47.4 1.2 29 75.1 43.1 34 53.6 447 1469 1979 18817 52.2 52.6 3968 1990 1980 12976 50.3 47.4 0.2 76 76.4 42.8 36.5 448 1468 12797 12986 50.2 40.9 36.9 36.9 36.9 36.9 36.9 3						1						0.5	280	74.8	42.	34	52.6
1451 1451 8867 8866 50.7 50 9861 9252 9255 50.1 50 0.6 366 76.4 44.8 34 54.1 432 1452 28939 28960 54.7 50 9862 23901 22922 55.3 55 0.6 366 76.6 45.2 34 55.1 433 1453 28940 28960 52.4 47.6 3985 28260 53.5 55.6 0.7 363 76.6 45.2 34 54.1 435 1454 28941 28960 50.9 45 3854 28298 28280 51.4 52.6 0.5 358 76.3 44.7 34 53.4 435 1455 3380 3380 51.4 42.9 3955 3494 3473 50.4 40.9 1 1355 73.7 45.9 34 51.8 437 1457 12373 12391 50.8 47.4 3957 12394 12376 50.3 47.4 0.4 622 76.4 42.9 34.5 437 1457 12373 12391 50.8 47.4 3957 12394 12376 50.3 47.4 0.4 622 76.4 42.9 34.5 439 1459 3361 3361 50.5 42.9 3958 19921 12376 50.3 47.4 0.4 622 76.4 42.9 34.5 440 1460 12234 12252 50.6 47.4 3961 12994 12376 50.3 47.4 0.2 76.5 42.6 34.5 53.8 441 1461 12234 12252 50.6 47.4 3961 12994 12376 50.3 47.4 0.2 76.5 76.5 42.6 34.5 53.8 442 1462 3034 3053 50.3 50.3 50.3 3962 3847 3828 50.6 47.4 0.2 76.7 76.5 42.8 34.5 53.8 445 1467 3034 3053 50.3 50.3 50.3 3962 3847 3828 50.6 47.4 0.2 76.7 76.4 42.2 34.5 53.6 445 1468 12274 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 76.7 76.5 76.5 42.6 34.5 53.8 446 1469 9926 9944 50.5 52.6 3966 16065 16065 16068 16											55	0.1	363	76.6	45.2	2 34	55.3
432 1452 28933 28960 54.7 50 3952 28301 28262 55.3 55.0 6.3 83 6.6 45.2 34 55.1 433 1453 28940 28960 52.4 47.6 3953 28900 28288 53.5 52.6 1 367 76.5 45 34 55.4 435 1455 3380 3380 51.4 42.9 3955 3694 3473 50.4 40.9 1 135 73.7 45.9 34 51.8 435 1455 13709 19730 51.3 40.9 3956 19916 18985 50.2 40.9 1 120 73.6 41.3 34 51.7 437 1457 12373 12391 50.8 47.4 3957 12294 12276 50.3 47.4 40.9 1 135 73.7 45.9 34 51.8 439 1459 3961 3381 3381 50.5 42.9 3895 3894 3473 50.4 40.9 1 126 72.6 43.8 34 51.7 439 1459 3361 3381 50.5 42.9 3895 3894 3473 50.4 40.9 0.1 134 73.8 40.3 34 51.7 440 1460 12234 12252 50.6 47.4 3960 12392 12376 50.3 47.4 0.9 0.1 134 73.8 40.3 34 51.7 441 1461 12234 12252 50.6 47.4 3961 12594 12376 50.3 47.4 0.2 761 76.4 42.2 34 53.7 443 1465 12224 12252 50.6 47.4 3961 12594 12576 50.3 47.4 0.3 61.1 76.4 42.2 34 53.7 443 1465 12224 12252 50.6 47.4 3961 12594 12576 50.3 47.4 0.2 761 76.4 42.2 34 53.4 443 1466 12224 12252 50.6 47.4 3965 12594 12576 50.3 47.4 0.2 761 76.4 42.2 34 53.6 444 1464 12726 12746 51.3 47.6 3964 12594 12576 50.3 47.4 1.7 388 75.1 41.2 34 52.8 445 1465 12234 12252 50.6 47.4 3965 12594 12576 50.3 47.4 1.7 388 75.1 41.2 34 52.8 446 1466 9929 9944 50.5 52.6 3966 10605 10568 51.1 50 0.6 680 75.8 41.2 34 53.6 447 1467 3034 3035 50.3 50.3 50.9 60 10605 10568 51.1 50 0.6 680 75.8 41.2 34 53.6 448 1468 12577 12586 50.2 40.0 3967 3968 13545 13527 50.3 50.6 60.0 60.0 76.5 76.5 76.4 42.5 34 53.6 449 1469 1477 18667						-					52.6	1.8	366	76.4	44.8	34	
432 1452 28983 28960 54.7 50 3962 29301 29262 55.3 55.6 0.6 9363 76.6 45.2 34 54.4 433 1453 28941 28960 52.4 47.6 3955 29280 29280 51.4 52.6 0.5 358 76.3 44.7 34 52.4 435 1455 3380 3380 51.4 42.9 3955 3494 3473 50.4 40.9 1 1355 73.7 45.9 34 51.8 437 1457 12373 12391 50.6 50.8 37.4 3957 12394 12376 50.2 40.9 1 1355 73.7 45.9 34 51.8 438 1458 19794 19813 50.5 50.8 47.4 3957 12394 12376 50.3 47.4 0.4 622 76.4 42.9 34.5 51.7 439 1459 3361 3381 50.5 42.9 3959 3494 3473 50.4 40.9 1 134 73.8 34.8 50.9 440 1460 12234 12252 50.6 47.4 3967 12994 12976 50.3 47.4 0.4 622 76.4 42.9 34.5 50.4 441 1461 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 76.5 42.6 34.5 53.8 442 1462 3034 3053 50.3 50.3 50.3 50.3 50.3 50.3 40.9 12976 50.3 47.4 0.2 76.5 42.6 34.5 53.8 445 1463 80867 8887 52.3 47.6 3962 3867 3862 50.6 47.4 0.2 76.5 76.5 42.6 34.5 53.8 445 1464 12726 12746 51.3 47.6 3968 12994 12976 50.3 47.4 0.2 76.7 76.4 42.2 34.5 52.8 445 1465 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 76.7 76.4 42.2 34.5 53.6 445 1465 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 76.7 76.4 42.8 34.5 53.8 445 1467 3034 3053 50.3									9235	50.1	50	0.6	386	75.1	41.5	34	52.7
433 1483 28940 28990 52.4 47.6 3953 29300 29288 53.5 52.6 1 367 76.5 45 34 54.4 434 1484 28941 28990 50.9 48.5 3956 3494 3473 50.4 40.9 1 135 73.7 45.9 34 51.8 435 1456 19709 19730 51.3 40.9 3956 19916 19895 50.2 40.9 1 208 73.6 41.3 34 51.7 437 1457 12373 12373 12373 12391 50.8 47.4 3957 12994 12376 50.3 47.4 0.4 622 76.4 42.9 34 53.7 438 1458 19794 19813 50 50 3958 19921 19900 51.8 45.5 1.8 122 72.6 43.8 34 50.9 440 1460 12224 12252 50.6 47.4 3957 12994 12376 50.3 47.4 0.4 622 76.4 42.9 34 53.7 441 1461 12224 12252 50.6 47.4 3950 12994 12376 50.3 47.4 0.4 622 76.4 42.9 34 53.7 441 1461 12224 12252 50.6 47.4 3950 12994 12376 50.3 47.4 0.2 761 76.4 42.4 34 53.7 442 1462 3034 3035 50.3 50.3 50.8 50.962 3847 3826 50.6 47.6 3963 3254 4224 4225 412252 50.6 47.6 3963 3254 4257 50.3 47.4 0.2 761 76.4 42.8 34 53.6 4441 1464 12726 12746 51.3 47.6 3963 3254 9226 50.6 47.4 1.7 3867 54.4 42.8 43.4 4	_								29282	55.3	55	0.6	363	76.6	45.2	34	
435 1455 3296 3390 3390 51.4 42.9 3955 3494 3475 50.4 40.9 11 325 73.6 437 34 51.8 436 1456 19709 19730 51.3 40.9 3956 19916 19895 50.2 40.9 11 208 73.6 41.3 34 51.7 437 1457 12373 12391 50.8 47.4 3957 12394 12376 50.3 47.4 0.4 622 76.4 42.9 34 53.7 438 1458 19794 19913 50.5 50.3 58.9 3494 3473 50.4 40.9 0.1 134 73.8 43.8 34 50.9 439 1459 3361 3381 50.5 42.9 3959 3494 3473 50.4 40.9 0.1 134 73.8 46.3 34 50.9 4401 1461 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.9 0.1 134 73.8 46.3 34 51.9 4421 462 3034 3053 50.3 5								29306	29288	53.5	52.6	1	367	76.5	45	34	
439 1456 1379 19730 51.3 40.9 3956 19916 1995 50.2 40.9 1 1 208 73.6 41.3 45 51.8 439 1456 1979 19730 51.3 40.9 3956 19916 1995 50.2 40.9 1 208 73.6 41.3 45 51.8 439 1457 1457 12373 12391 50.8 47.4 3957 12994 12976 50.3 47.4 0.4 622 76.4 42.9 43 53.7 439 1458 19794 19813 50 50 3958 19921 19900 51.8 45.5 1.8 128 72.6 43.8 34 55.7 44 14 61 1461 12224 12225 50.6 47.4 3960 12992 12974 51.2 52.6 0.6 75 76.5 42.6 43.8 45.5 440 1460 12224 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 761 76.4 42.4 34 53.7 442 1462 3934 3053 50.3 50.3 50.3 50.3 50.3 50.3 50.3								29298	29280	51.4	52.6	0.5	358	76.3	44.7	34	
439 1496 1970 19730 51.3 40.9 3986 19916 19985 50.2 40.9 1 208 73.6 41.3 34 51.7 439 1458 19794 19813 50 50 3958 19921 19900 51.8 45.5 1.8 128 72.6 43.8 34 50.9 439 1459 3361 3381 50.5 42.9 3959 3494 3473 50.4 40.9 0.1 134 73.8 46.3 34 51.9 440 1460 12234 12252 50.6 47.4 3961 12992 12974 51.2 52.6 6.6 75.9 76.5 42.6 345 53.4 441 1461 12234 12252 50.6 47.4 3961 12992 12976 50.3 47.4 0.2 761 76.4 42.4 45 53.6 442 1462 3034 3053 50.3 50 3962 3347 3628 50.6 47.6 36.3 47.6 36.3 442 1462 3034 3053 50.3 50 3962 3347 3628 50.6 47.6 36.3 47.6 36.3 444 1464 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1 299 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3966 12998 12979 50.1 45.0 5.7 765 76.4 42.5 34 445 1467 3034 3053 50.3 50 3967 3846 3625 52 40.9 1.7 613 76.3 42.7 34 53.6 447 1467 3034 3053 50.3 50 3967 3846 3625 52 40.9 1.7 613 76.3 42.7 34 53.6 448 1468 12977 12986 50.2 40 3968 13545 13527 50.3 52.6 0.5 699 77.4 45.5 44.5 44.9 14.9 44.9 14.9 14.9 44.9 44.9 14.9 44.									3473	50.4	40.9	1	135	73.7	45.9	34	
439 1489 19794 19813 50 60 3988 19921 19906 50.3 47.4 0.4 622 76.4 42.9 34 53.7 439 1489 3361 3381 50.5 42.9 3859 3494 3473 50.4 40.9 0.1 134 73.8 45.3 45.9 440 1460 12234 12252 50.6 47.4 3960 12992 12974 51.2 52.6 0.6 75.9 76.5 42.6 34 53.8 441 1461 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.6 0.3 61.4 442 1462 3034 3063 50.3 50.3 50.3 50.3 62.8 60.6 60.3 60.4 60.4 60.4 443 1464 14726 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 1.7 388 75.1 41.2 34 53.8 443 1463 8867 8867 52.3 47.6 3964 12994 12976 50.3 47.4 1.7 388 75.1 41.2 34 52.8 444 1464 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1.7 388 75.1 41.2 34 52.8 445 1465 12234 12252 50.6 47.4 3965 12998 12979 50.1 45 50.5 76.4 42.5 43 53.6 446 1466 9926 9944 50.5 52.6 3966 10605 10588 51.1 50 0.6 680 75.6 41.2 34 53.3 448 1468 12977 12996 50.2 40 3968 13545 13527 50.3 52.6 0 569 77.4 45.5 34 53.4 449 1469 19799 19817 52.2 52.6 3966 19909 19865 52.5 40 0.3 1111 71.6 43.2 34 53.8 450 1470 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 300 75.1 41.1 34 52.7 451 1471 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 300 75.7 42.5 53.7 452 1472 10141 10160 51 45.3 372 10608 10589 51 50 0.3 47.6 40.4 30.5 52.8 453 1473 3192 3213 51.8 45.5 3975 3494 3473 50.4 40.9 1.4 30.3 74.9 41.9 35 52.8 454 1476 20750 20774 51.6 40.9 3976 10605 10589 51 50 0.3 47.6 40.4 35 52.4 459 1479 8067 8067 8067 8068 8068 80.6 44.5 50.0 3.6 47.4 40.2 35 53.1 450 1476 20750 20774 51.6 40						_			19895	50.2	40.9	1	208	73.6	41.3	34	
439 1459 3361 3361 505 503 503 505 3458 19921 19900 51.8 45.5 1.8 1.28 72.6 43.8 34 50.9 440 1460 12234 12252 50.6 47.4 3660 12992 12974 51.2 52.6 0.6 759 76.5 42.6 34 53.8 441 1461 12234 12252 50.6 47.4 3660 12992 12974 51.2 52.6 0.6 759 76.5 42.6 34 53.8 441 1461 12234 12252 50.6 47.4 3660 12992 12976 50.3 47.4 0.2 761 76.4 42.4 34 53.8 442 1462 3034 3035 50.3 50 362 3647 3628 50.6 47.4 1.7 368 36 75.1 41.2 34 52.8 444 1464 12726 12746 51.3 47.6 3663 9254 9236 50.6 47.4 1.7 368 75.1 41.2 34 52.8 444 1464 12726 12746 51.3 47.6 3664 12994 12976 50.3 47.4 1.7 269 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3665 12998 12979 50.1 45 50.5 765 76.4 42.5 34 53.8 447 1467 3034 3033 50.3 50.3 60.6 60.66 10.665 10.688 51.1 50 0.6 66.0 76.8 41.2 41.5 449 1469 12977 12966 50.2 40.3 368 13.545 13.527 50.3 52.6 0.5 699 77.4 45.5 45.5 449 1469 19799 19817 52.2 52.6 3668 19909 19865 52.5 40.9 1.7 61.3 76.3 42.7 34 53.6 450 1470 8867 8887 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 380 75 41.1 34 52.7 452 1472 10141 10180 51 45 3972 10608 10589 51 50 0.3 47.6 74.9 40.2 35 53.7 452 1472 10141 10180 51 45 3972 10608 10589 51 50 0.4 68 74.9 40.2 35 52.6 456 1476 10250 10274 51.6 52.6 3978 27566 27546 50.7 47.6 0.7 20.0 74.8 44.3 35 52.6 457 1477 27367 27385 51.4 52.6 3978 27576 27556 51 40.9 0.5 53.6 74.6 40.4 35 52.6 458 1478 27367 27385 51.4 52.6 3989 27576 27556 51 40.9 0.5 51.2 75.5 41.2 35 53.4 459 1488 18794 18724 50.8 47.6 39879 3936 39						47.4	3957	12994	12976	50.3	47.4	0.4	622	76.4	42.9	34	
449 1469 3361 3381 50.5 42.9 3959 3494 3473 50.4 40.9 0.1 134 73.8 46.3 34 51.9 440 1460 12234 12252 50.6 47.4 3960 12992 12974 51.2 52.6 0.6 75.9 76.5 42.6 34 53.6 442 1462 3034 3053 50.3 50.3 50.3 50.2 3662 3347 3628 50.6 47.6 0.2 761 76.4 42.4 34 53.7 442 1462 3034 3053 50.3 50.3 50.3 50.2 3622 3347 3628 50.6 47.6 0.3 614 76.4 42.8 34 53.6 444 1464 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1 229 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3966 12998 12979 50.1 45 0.5 765 76.4 42.5 34 53.6 445 1466 9926 9926 9944 50.5 52.6 3968 12998 12979 50.1 45 0.5 765 76.4 42.5 34 53.6 447 1467 3034 3053 50.3 50 3867 3646 3625 52 40.9 1.7 613 76.3 42.7 34 53.6 448 1468 12977 12986 50.2 40 3968 13545 13527 50.3 52.6 0.5 569 77.4 45.5 34 53.3 449 1469 19799 1917 52.2 52.6 3969 13985 52.5 40.9 1.7 613 76.3 42.7 34 53.6 450 1470 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 330 75 41.1 34 52.7 452 1472 10141 10160 51 45 3972 10608 10698 51.5 50 0.3 47.6 57.7 42.3 53.7 455 1473 3192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 30.3 74.9 41.9 35 52.6 456 1476 10250 10274 51.6 40 3976 10608 10588 51.1 50 0.5 56 74.6 40.4 35 52.6 457 1477 27367 27385 51.4 52.6 3979 376 9355 51 40.9 1.3 510 75.7 41.8 35 52.6 458 1473 3308 379	_							19921	19900	51.8	45.5	1.8	128	72.6	43.8		
440 1460 12234 12252 50.6 47.4 3960 12994 12976 50.3 47.4 30.6 76.5 42.6 34 53.8 441 1461 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 761 76.4 42.8 34 53.8 442 1462 3034 3063 50.3 50 3662 3647 3628 50.6 47.4 1.7 388 75.1 41.2 34 53.4 443 1464 14726 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1.7 388 75.1 41.2 34 52.8 444 1464 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1 299 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3965 12998 12979 50.1 45 50.5 76.6 42.5 34 53.6 446 1466 9926 9944 50.5 52.6 3966 10605 10588 51.1 50 0.6 680 75.6 41.2 34 53.3 448 1468 12977 12996 50.2 40 3688 13645 13527 50.3 52.6 0 569 77.4 45.5 34 53.4 449 1469 19799 19817 52.2 52.6 3969 19909 19865 52.5 40 0.3 1111 71.6 43.2 34 50.8 450 1470 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 300 75.1 41.1 34 52.7 452 1472 10141 10160 51 45 3072 10608 10589 51 50 0.3 476 74.9 40.2 35 53.8 453 1473 33192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 30.3 74.9 41.9 35 52.6 456 1476 20250 10274 51.6 40 3976 10605 10589 51 50 0.3 476 40.4 35 52.7 457 1477 27367 27385 51.4 52.6 3975 27566 27546 50.7 47.6 0.7 200 74.8 44.3 35 52.7 458 1478 27367 27385 51.4 52.6 3978 27571 27551 51.4 42.9 0.25 74.6 43.3 35 52.7 459 1479 8867 8867 8867 8868 8								3494	3473	50.4	40.9	0.1	134	73.8		-	
441 1461 12234 12252 50.6 47.4 3961 12994 12976 50.3 47.4 0.2 761 76.4 42.4 34 53.7 442 1462 3034 3035 50.3 50.3 3628 3867 3628 50.6 45 0.3 614 76.4 42.8 34 53.7 443 1463 8867 8867 52.3 47.6 3663 9254 9236 50.6 47.4 1.7 386 75.1 41.2 34 52.8 444 1464 12726 12746 51.3 47.6 3664 12994 12976 50.3 47.4 1 269 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3666 12998 12979 50.1 45 0.5 765 76.4 42.5 34 53.6 446 1466 9926 9944 50.5 52.6 3966 10605 10588 51.1 50 0.6 660 76.8 41.2 34 53.3 447 1467 3034 3053 50.3 50.3 667 3646 3625 52 40.9 1.7 613 76.3 42.7 34 53.6 449 1469 19799 19817 52.2 52.6 3968 19909 19865 52.5 40.9 1.7 613 76.3 42.7 34 53.6 449 1469 19799 19817 52.2 52.6 3968 19909 19865 52.5 40.9 1.7 613 76.3 42.7 34 53.6 450 1470 8867 8887 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 380 75 41.1 34 52.7 452 1472 10141 10180 51 45 3972 10608 10589 51 50 0.4 48 74.9 40.2 35 53.7 452 1472 10141 10180 51 45 3972 10608 10589 51 50 0.4 48 74.9 40.2 35 52.6 456 1476 3972 27385 51.4 52.6 3978 27566 27546 50.7 47.6 0.7 20.7 74.8 44.3 36.8 37.5 37.8 34.9 37.8		1				47.4	3960	12992	12974	51.2	52.6	0.6	759	76.5			
442 1462 3934 3953 50.3 50.3 50.3 50.3 50.3 50.3 50.2 50.6 47.4 1.7 348 75.1 42.8 34 53.6 444 1464 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1 299 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3965 12998 12979 50.1 45 0.5 765 76.4 42.5 34 53.6 446 1466 9926 9926 9944 50.5 52.6 3966 12998 12979 50.1 45 0.5 765 76.4 42.5 34 53.6 447 1467 3034 3053 50.3 50 3667 3646 3625 52 40.9 1.7 613 76.3 42.7 34 53.6 448 1468 12977 12996 50.2 40 3968 13545 13527 50.3 52.6 0 569 77.4 45.5 34 53.6 449 1469 19799 1917 52.2 52.6 3969 19909 19865 52.5 40 0.3 111 71.6 34.2 34 53.6 450 1470 8867 8867 52.3 47.6 3971 9342 9323 52.1 50 0.3 47.6 75.7 42 35 53.7 451 1471 8867 6887 52.3 47.6 3971 9342 9323 52.1 50 0.3 47.6 75.7 42 35 53.7 452 1472 10141 10160 51 45 3972 10608 10589 51 50 0.3 47.6 57.5 42 35 53.7 453 1473 3192 3213 51.6 45.5 3973 3494 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 456 1476 10250 10274 51.6 40 3976 10608 10588 51.1 50 0.5 356 74.6 44.1 35 52.4 457 1477 27367 27335 51.4 52.6 3979 27568 27564 50.7 40.9 1.3 51.0 75.7 41.8 35 52.6 458 1478 27367 27335 51.4 52.6 3977 27568 27564 50.7 40.9 1.3 51.0 75.7 41.8 35 52.6 450 1480 1481 148		_						12994	12976	50.3	47.4	0.2	761	76.4	42.4	34	
443 1464 14726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1.7 3.86 75.1 41.2 34 52.8 444 1464 14726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1 299 75.1 41.2 34 53.6 446 1466 9926 9944 50.5 52.6 3966 10905 10588 51.1 50 0.6 680 75.8 41.2 34 53.4 447 1467 3034 3053 50.3 50 3967 3646 3625 52 40.9 1.7 61.3 76.3 42.7 34 53.6 448 1488 12977 12996 50.2 40 3968 13945 13527 50.3 52.6 0 569 77.4 45.5 34 53.4 449 1469 19799 19817 52.2 52.6 3969 19900 19865 52.5 40 0.3 111 71.6 43.2 34 53.8 450 1470 8867 8887 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 390 75 41.1 34 52.7 451 1471 8867 8887 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 390 75 41.1 34 52.7 452 1472 10141 10160 51 45 3972 10608 10589 51 50 0 468 74.9 40.2 35 53.8 452 1472 10141 10160 51 45 3972 10608 10589 51 50 0 468 74.9 40.2 35 52.8 453 1473 33192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 30.3 74.9 41.9 35 52.6 456 1476 27367 27385 51.4 52.6 3975 27566 27546 50.7 47.6 0.7 200 74.8 44.1 35 52.7 459 1479 8867 8867 52.3 47.6 3979 3976 10605 10588 51.1 50 0.5 36.6 74.6 40.4 35 52.7 459 1479 8867 8868 8867 52.3 47.6 3979 3976 10605 10588 51.1 40.9 0.3 21.3 75.4 44.3 35 52.7 450 1479 27367 27385 51.4 52.6 3978 27579 27558 51.4 40.9 0.3 21.3 75.4 44.3 35 52.4 450 1479 3868 8867 8867 8868 8867 8867 8868 8867 8867 8868 8867 8868 886							3962	3647	3628	50.6	45	0.3	614	76.4	42.8		
444 1464 12726 12746 51.3 47.6 3964 12994 12976 50.3 47.4 1 269 75.1 43.1 34 52.8 445 1465 12234 12252 50.6 47.4 3965 12998 12979 50.1 45 0.5 765 76.4 42.5 34 53.3 447 1467 3034 3053 50.3 50.3 50.3 60.6 56.6 50.6 66.0 75.8 41.2 34 53.3 447 1467 3034 3053 50.3		1.100				47.6	3963	9254	9236	50.6	47.4	1.7	388	75.1	41.2		
446 1466 9926 9926 9944 50.5 52.6 3966 12998 12999 50.1 45 0.5 765 76.4 42.5 34 53.6 447 1467 3034 3053 50.3 50 3967 3646 3625 52 40.9 1.7 613 76.3 42.7 34 53.6 448 1468 12977 12996 50.2 40 3968 13545 13527 50.3 52.6 0 569 77.4 45.5 34 53.8 449 1469 19799 19187 52.2 52.6 3969 19909 19885 52.5 40 0.3 111 71.6 34.2 34 53.4 450 1470 8867 8867 52.3 47.6 3971 9342 9323 52.1 50 0.3 47.6 75.7 42 35 53.7 451 1471 8867 8867 52.3 47.6 3971 9342 9323 52.1 50 0.3 47.6 75.7 42 35 53.7 452 1472 10141 10160 51 45 3972 10608 10589 51 50 0.3 47.6 37.7 40.2 35 52.5 453 1473 3192 3213 51.8 45.5 3973 3484 3473 50.4 40.9 1.4 303 74.9 40.2 35 52.6 456 1474 3380 3379 50.7 45 3974 3647 3628 50.6 45 0.1 288 75.5 43.8 35 53.1 457 1477 27367 27395 51.4 52.6 3976 10605 10688 51.1 50 0.5 356 74.6 40.4 35 52.6 458 1476 10250 10274 51.6 40 3976 10605 10688 51.1 50 0.5 356 74.6 40.4 35 52.6 459 1477 27367 27385 51.4 52.6 3977 27568 27546 50.7 40.9 1.3 510 75.7 41.8 35.5 52.6 459 1479 8867 8867 52.3 47.6 3979 3376 9355 51 40.9 0.3 213 75.5 41.2 35 53.4 450 1480 27367 27385 51.4 52.6 3980 27579 27558 51.4 40.9 0.3 213 75.5 41.2 35 53.4 460 1480 27367 27385 51.4 52.6 3980 27579 27558 51.1 40.9 0.3 213 75.5 41.2 35 53.4 461 1481 27367 27385 51.4 52.6 3980 27579 27558 51.1 40.9 0.3 213 75.5 41.2 35 53.4 462 1482 18704 18724 50.8 47.6 3983 19217 19196 50.2 40.9 0.5 512 75.5 41.2 35 53.4 463 1483 18704 18724 50.8 47.6 3988 3980 37579 27558 51.1 40.9 0.3 2								12994	12976	50.3	47.4	1	269	75.1	43.1	34	
446 1466 9926 9944 50.5 52.6 3966 10605 10588 51.1 50 0.6 680 75.8 41.2 34 53.3 447 1467 3034 3053 50.3 50 3967 3646 3625 52 40.9 1.7 613 76.3 42.7 34 53.3 448 1468 12977 12986 50.2 40 3968 13945 13527 50.3 52.6 0 569 77.4 45.5 34 54.3 449 1469 19799 19817 52.2 52.6 3969 19909 19865 52.5 40 0.3 111 71.6 43.2 34 50.8 450 1470 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 390 75 41.1 34 52.7 451 1471 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 390 75 41.1 34 52.7 452 1472 10141 10160 51 45 3972 10606 10589 51 50 0 468 74.9 40.2 35 53.7 452 1472 10141 10160 51 45 3973 3494 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 454 1474 3360 3379 50.7 45 3974 3467 3628 50.6 45 0.1 286 55.5 55.5 43.6 455 1476 10250 10274 51.6 40 3976 10605 10589 51.1 50 0.5 366 74.6 40.4 35 52.7 459 1479 8867 8867 52.3 47.6 3978 27561 27551 51.4 42.9 0.205 74.6 40.4 35 52.4 450 1479 8867 8867 52.3 47.6 3979 9376 9355 51 40.9 1.3 510 75.7 41.8 35 52.4 460 1480 27367 27385 51.4 52.6 3978 27576 27555 51.4 40.9 0.3 21.3 75 44.6 35 52.4 451 1471 27367 27385 51.4 52.6 3978 27576 27555 51.4 40.9 0.3 21.3 75 44.6 35 52.4 451 1481 27367 27385 51.4 52.6 3980 27576 27555 51.4 40.9 0.3 21.3 75 44.6 35 52.4 462 1482 1870 1872 47.6 30.8 47.6 3989 27576 27555 51.4 40.9 0.3 21.3 75 44.6 35 52.9 462 1482 1870 1871 1872 50.8 47.6 3980 3980 3115 51.9 40.9 0.5 51.4 75.5 41.2 35 53.1 463 1483 3361 3361 50.5 42.9 3987 3646 3625 52 40.9 1.5 526 75.6 41.3 35 53.1							3965	12998	12979	50.1	45	0.5	765	76.4			
447 1467 3034 3053 50.3 50.3 50.3 50.3 50.3 50.3 50.4 3068 35.5 52 40.9 1.7 61.3 76.3 42.7 34 53.6 449 1469 19799 19817 52.2 52.6 3068 19909 19805 52.5 40 0.3 111 71.6 43.2 45 53.4 449 1469 19799 19817 52.2 52.6 3068 19909 19805 52.5 40 0.3 111 71.6 43.2 45 50.8 450 1470 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 380 75 41.1 34 52.7 451 1471 10160 51 45 3972 10608 10589 51 50 0.3 476 75.7 42 35 53.7 452 1472 10141 10180 51 45 3972 10608 10589 51 50 0.3 476 75.7 42 35 53.7 455 1473 3192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 455 1475 27367 27385 51.4 52.6 3978 27566 27546 50.7 47.6 60.7 20.7 74.8 44.5 55 52.7 455 1475 10200 10274 51.6 40 3976 10605 10588 51.1 50 0.5 356 74.6 40.4 35 52.6 459 1478 27367 27385 51.4 52.6 3978 27576 27556 27546 50.7 40.9 1.2 20.2 74.6 44.1 35 52.6 459 1478 27367 27385 51.4 52.6 3977 27568 27548 50.7 40.9 1.2 20.2 74.6 44.1 35 52.6 459 1479 8867 82.3 47.6 3979 27571 27551 51.4 42.9 0.2 20.7 44.6 43.9 55.2 42.9 1.7 27367 27385 51.4 52.6 3987 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.4 461 1481 27367 27385 51.4 52.6 3981 27576 27555 51 40.9 0.3 213 75.5 41.2 35 53.4 465 1481 27367 27385 51.4 52.6 3981 27579 27558 51.1 40.9 0.3 213 75.4 40.9 35 52.6 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 51.2 75.5 41.2 35 53.4 465 1485 27365 27384 52.6 50.3986 13155 13137 50.4 40.9 1.5 52.0 75.6 41.3 35 53.1 465 1485 27365 27384 52.6 50.3986 13155 13137 50.4 4					50.5	52.6	3966	10605	10588	51.1	50	0.6	680	75.8			
449 1469 19799 19817 52.2 40 3968 13545 13527 50.3 52.6 0 569 77.4 45.5 34 54.3 449 1469 18799 19817 52.2 52.6 3669 19909 19885 52.5 40 0.3 111 71.6 32.2 34 53.3 45.1 34 52.7 45.1 34 52.7 45.1 34 52.7 34.5 34						50	3967	3646	3625	52	40.9	1.7	613				
449 1469 19799 19817 52.2 52.6 3969 19909 1985 52.5 40 0.3 111 71.6 43.2 34 50.8 450 1471 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 380 75 41.1 34 52.7 451 1471 8867 8867 52.3 47.6 3970 9246 9226 50.5 42.9 1.8 380 75 41.1 34 52.7 34.5			-			40	3968	13545	13527	50.3	52.6	0					
450 1470 8867 8867 52.3 47.6 3971 9246 9226 50.5 42.9 1.8 380 75 41.1 34 52.7 451 1471 8867 8887 52.3 47.6 3971 9342 9323 52.1 50 0.3 476 75.7 41.3 34 52.7 42 35 53.4 452 1472 10141 10180 51 45 3972 10608 10589 51 50 0 486 74.9 40.2 35 52.8 453 1473 3192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 454 1474 3360 3379 50.7 45 3974 3647 3628 50.6 45 0.1 288 .75.5 41.9 35 52.6 455 1475 27367 27385 51.4 52.6 3975 27566 27546 50.7 47.6 0.7 200 74.8 44.5 35 52.7 45 1475 27367 27385 51.4 52.6 3976 27568 27548 50.2 42.9 1.2 202 74.6 40.4 35 52.6 459 1478 27367 27385 51.4 52.6 3976 27568 27548 50.2 42.9 1.2 202 74.6 40.4 35 52.6 459 1479 8867 52.3 47.6 3979 9376 27558 51.4 42.9 0 205 74.6 40.4 35 52.4 459 1479 8867 8867 52.3 47.6 3979 9376 9355 51 40.9 1.3 510 75.7 41.8 35 53.4 460 1480 27367 27385 51.4 52.6 3980 27576 27556 51.4 40.9 1.3 510 75.7 41.8 35 53.4 461 1481 27367 27385 51.4 52.6 3980 27576 27556 51.4 40.9 0.3 213 57.5 44.3 35 52.8 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.4 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.4 461 1484 18686 18715 51.7 50 3985 19217 19196 50.2 40.9 1.5 520 75.6 41.3 35 53.4 465 1485 27365 27384 52.6 50 3985 27464 27443 54.4 50.5 52.0 40.9 1.5 520 75.6 41.3 35 53.1 465 1485 27365 27384 52.6 50 3985 27464 27443 54.4 50.5 52.9 57.6 41.3 35 53.1 465 1485 27365 27384 52.6 50 3985 27464 27443 54.4 55.5 1.4 10.0 70.8 43 35 50.4 461 1481 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 520 75.6 41.3 35 53.1 468 1488 3361 3361 50.5 42.9 3987 3646 3625 52 40.9 1.5 520 75.6 41.3 35 53.1 468 1488 3361 3361 50.5 42.9 3987 3646 3625 52 40.9 1.5 520 75.6 41.3 35 53.1 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 664 75.5 40.5 35 53.1 471 1491 3782 3801 51.3 50 3991 3155 313774 3777 3777 50.4 40.9 521 75.6 44.0 35 53.1 471 1491 3782 3801 31.3 50 3991 3155 31377 3777 3777 3778 50.0 40.9 50.0 50.0 50.0 50.0 50.0 50.0 50.0 5						52.6	3969	19909	19885	52.5	40	0.3	111				
451 1471 8867 8867 8867 82.3 47.619371 9342 9323 52.1 50 0.3 476 75.7 42 35 53.7 452 1473 3192 3213 51.8 45.53973 3484 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 453 1473 3192 3213 51.8 45.53973 3484 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 455 1474 3360 3379 50.7 45 3974 3647 3628 50.6 45 0.1 288 75.5 43.8 35 53.1 455 1475 27367 27365 51.4 52.6 3975 27566 27546 50.7 47.6 0.1 288 75.5 43.8 35 53.1 457 1477 27367 27365 51.4 52.6 3977 27568 27546 50.7 47.6 0.1 288 75.5 43.8 35 52.6 451 1477 27367 27365 51.4 52.6 3977 27568 27546 50.7 47.6 0.5 356 74.6 40.4 35 52.6 458 1478 27367 27365 51.4 52.6 3977 27568 27546 50.2 42.9 1.2 20.2 74.6 44.1 35 52.4 458 1478 27367 27365 51.4 52.6 3979 2376 9355 51 40.9 1.3 510 75.7 41.8 35 53.4 450 1480 27367 27365 51.4 52.6 3980 27576 27555 51 40.9 1.3 510 75.7 41.8 35 53.4 451 1481 27367 27365 51.4 52.6 3980 27576 27558 51.1 40.9 0.4 210 74.8 44.3 35 52.8 451 1482 13704 13724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 445 1483 13704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 445 1485 1488 1488 18866 18715 51.7 50 3984 19217 19196 50.2 40.9 0.5 512 75.5 41.2 35 53 465 1485 27367 27384 52.6 50 3985 27464 27443 54 55.5 1.4 10.0 70.8 43 35 50.4 466 1486 18866 18715 51.7 50 3986 19217 19194 50.2 40.9 0.5 512 75.6 41.3 35 53.4 466 1486 18866 18715 51.7 50 3986 19217 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 465 1485 27365 27384 52.6 50 3985 27464 27443 54 45.5 1.4 10.0 70.8 43 35 50.4 466 1486 18866 18715 51.7 50 3986 19217 19194 50.2 40.9 1.5 520 75.6 41.4 35 53.1 468 1488 3361 3381 50.5 42.9 3988 3647 3628 50.6 44.5 0.1 287 75.6 43.3 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 526 75.6 43.3 35 53.1 469 1489 3782 3801 51.3 50 3989 34445 4425 50.6 42.9 0.7 663 75.5 43.3 35 53.1 469 1489 3782 3801 51.3 50 3989 3155 31375 51.5 50.0 44.4 4424 50.6 42.9 0.7 663 75.5 40.5 35 53 53 53 53 50 50 50 50 50 50 50 50 50 50 50 50 50					-	47.6	3970	9246	9226	50.5	42.9	1.8					
452 472 10141 10160 51 45 3972 10608 10589 51 50 0 468 74.9 40.2 35 52.8 453 473 3192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.8 454 474 3360 3379 50.7 45 3974 3647 3628 50.6 45 0.1 288 75.5 43.8 35 53.1 455 1475 27367 27365 51.4 52.6 3975 27566 27546 50.7 47.6 0.7 200 74.8 44.5 35 52.7 456 1476 10250 10274 51.6 40 3976 10605 10588 51.1 50 0.5 356 74.6 40.4 35 52.6 457 1477 27367 27365 51.4 52.6 3976 10605 10588 51.1 50 0.5 356 74.6 40.4 35 52.4 458 1478 27367 27385 51.4 52.6 3976 27568 27548 50.2 42.9 1.2 20.2 74.6 44.1 35 52.4 459 1479 8867 8887 52.3 47.6 3979 9376 9355 51 40.9 1.3 510 75.7 41.8 35 53.4 450 1480 27367 27385 51.4 52.6 3979 27576 27555 51 40.9 1.3 510 75.7 41.8 35 53.4 460 1480 27367 27385 51.4 52.6 3981 27579 27558 51.1 40.9 0.3 213 75 44.6 35 52.9 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 463 1483 18704 18724 50.8 47.6 3985 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 464 1484 18666 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 465 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 520 75.6 41.3 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.0 75.6 41.3 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.6 75.6 41.3 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.6 0.3 117 73.4 47 35 53.1 469 1489 3782 3301 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 3						47.6	3971	9342	9323	52.1	50	0.3	476	75.7			
483 1473 3192 3213 51.8 45.5 3973 3494 3473 50.4 40.9 1.4 303 74.9 41.9 35 52.6 485 1475 27367 27385 51.4 52.6 3975 27566 27546 50.7 47.6 0.7 200 74.8 44.5 35 52.7 486 1476 10250 10274 51.6 40 3976 10605 10588 51.1 50 0.5 35.6 74.6 40.4 35 52.6 487 1477 27367 27385 51.4 52.6 3977 27568 27548 50.2 42.9 1.2 202 74.6 40.4 35 52.6 488 1478 27367 27385 51.4 52.6 3977 27568 27548 50.2 42.9 1.2 202 74.6 40.4 35 52.6 489 1478 27367 27385 51.4 52.6 3978 27571 27551 51.4 42.9 0 205 74.6 43.9 35 52.7 489 1479 8867 8867 82.3 47.6 3979 9376 9355 51 40.9 0.3 21.3 510 75.7 41.8 35 53.4 461 1481 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.8 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 463 1484 18696 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 514 75.5 41.2 35 53 465 1485 27365 27384 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.8 43 35 53.1 466 1486 18696 18715 51.7 50 3986 19217 19196 50.2 40.9 1.5 520 75.6 41.3 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.4 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.0 75.6 43.9 35 53.1 470 1490 13039 13058 51.3 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 55 53.1 471 1491 3782 3801 51.3 50 3991 4444 4444 4444 4424 50.6 42.9 0.7 664 75.5 40.5 53.1 471 1490 9929 9946 50 50 3999 13155 13137 52.1 52.6 0.3 117 73.4 47 55 53.1 472 1499 9929 9946 50 50 3999 13155 13137 52.1 52.6 0.3 117								10608	10589	51	50	0	468	74.9			
455 1476 27367 27365 51.4 52.6 3975 27566 27546 50.7 47.6 0.7 200 74.8 44.5 35 53.1 456 1476 10250 10274 51.6 40 3976 10605 10588 51.1 50 0.5 368 74.6 40.4 35 52.4 457 1477 27367 27385 51.4 52.6 3977 27568 27548 50.2 42.9 1.2 202 74.6 44.1 35 52.4 458 1478 27367 27385 51.4 52.6 3978 27571 27551 51.4 42.9 0.2 205 74.6 44.1 35 52.4 459 1479 8867 8867 52.3 47.6 3978 27571 27551 51.4 42.9 0.2 205 74.6 43.9 35 52.4 460 1480 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.4 461 1481 27367 27385 51.4 52.6 3980 27576 27556 51 40.9 0.4 210 74.8 44.3 35 52.8 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.1 464 1484 18866 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 512 75.6 41.2 35 53.1 465 1486 18686 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 514 75.5 41.2 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.3 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.3 35 53.1 469 1489 3782 3801 51.3 50 3991 4445 4425 50.6 42.9 0.7 663 75.5 40.5 35.3 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52.1 471 1491 3782 3201 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 471 1499 9299 946 50 50 3992 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52.9 475 1499 9999 9946 50 50 3992 13157 13765 50.8 40.9 0.7 663 75.5 40.6 35 53.1 476 1494 9929 9946 50 50 3992 131747 13765 50.6 40.9 0.7 663 75.5 40.6 3			-					3494	3473	50.4	40.9	1.4	303	74.9			
486 1476 10250 10274 51.6 40.0 3976 10605 10588 51.1 50.0 74.6 0.7 20.0 74.8 44.5 35 52.7 486 1476 10250 10274 51.6 40.0 3976 10605 10588 51.1 50.0 5.366 74.6 0.41 35 52.4 487 1477 27367 27365 51.4 52.6 3977 27568 27548 50.2 42.9 1.2 20.2 74.6 44.1 35 52.4 489 1478 27367 27365 51.4 52.6 3978 27571 27551 51.4 42.9 0 20.5 74.6 43.9 35 52.7 480 1480 27367 27365 51.4 52.6 3979 9376 9355 51 40.9 1.3 510 75.7 41.8 35 53.4 480 1480 27367 27365 51.4 52.6 3980 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.8 481 1481 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.8 482 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.1 483 1483 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.1 485 1485 27365 27385 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.8 43 35 50.4 486 1486 18686 18715 51.7 50 3986 19217 19196 50.2 40.9 0.5 512 75.6 41.3 35 53.1 487 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.3 35 53.1 488 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 528 67.5 43.3 35 53.1 489 1489 3782 3801 51.3 50 3980 4445 4425 50.6 42.9 0.7 663 75.5 40.6 35 53.1 489 1489 3782 3801 51.3 50 3980 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52.4 481 1494 9929 9946 50 50 3992 13747 13726 50.6 42.9 0.7 663 75.5 40.6 35 53.1 489 1499 2929 9946 50 50 3992 13747 13726 50.6 40.9 0.7 663 75.5 40.6 35 53.1 489 1499 9929 9946 50 50 3992 13747 13726 50.6 40.9 0.7 663 75								3647	3628	50.6	45	0.1	288	.75.5	43.8		
486 14/6 10250 10274 51.6 40 3976 10605 10588 51.1 50 0.5 35.6 74.6 40.4 35 52.6 457 1477 27367 27385 51.4 52.6 3977 27568 27548 50.2 42.9 1.2 20.2 74.6 44.1 35 52.6 45.8 1478 27367 27385 51.4 52.6 3978 27571 27551 51.4 42.9 0 20.5 74.6 43.9 35 52.7 45.9								27566	27546	50.7	47.6	0.7	200	74.8	44.5		
489 1478 27387 27385 51.4 52.6 3977 27588 27548 50.2 42.9 1.2 20.2 74.6 44.1 35 52.4 458 1479 8867 27385 51.4 52.6 3978 27571 27551 51.4 42.9 0 20.5 74.6 43.9 35 52.7 459 1479 8867 8867 52.3 47.6 3979 9376 9356 51 40.9 1.3 510 75.7 41.8 35 53.4 460 1480 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.8 461 1481 27367 27385 51.4 52.6 3981 27579 27558 51.1 40.9 0.3 213 75 44.6 35 52.8 462 1482 18704 18724 50.8 47.6 3983 19217 19196 50.2 40.9 0.5 512 75.5 41.2 35 53.4 463 1483 18704 18724 50.8 47.6 3983 19217 19196 50.2 40.9 0.5 512 75.5 41.2 35 53.1 464 1484 18686 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 512 75.6 41.3 35 53.1 465 1485 27365 27365 27365 27334 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.8 43 35 50.4 466 1486 18696 18715 51.7 50 3986 19217 19196 50.2 40.9 1.5 520 75.6 41.3 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 286 75.5 43.7 35 53.1 469 1489 3782 3801 51.3 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52.4 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 664 75.5 40.6 35 53.1 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52.1 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 471 1491 3782 3801 51.3 50 3991 31747 31726 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.7 663 75.5 40.6 35 53.1 474 1494 9929 9946 50 50 3994 10439 10431 50.9 474 0.9 521		-		_				10605	10588	51.1	50	0.5	356	74.6	40.4		
489 1479 8867 8867 52.3 47.6 3979 27551 51.4 42.9 0 205 74.6 43.9 35 52.7 489 1479 8867 8887 52.3 47.6 3979 3976 39356 51.4 40.9 1.3 510 75.7 41.8 35 53.4 480 1480 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 21.0 74.8 44.3 35 52.8 481 1481 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 21.0 74.8 44.3 35 52.8 482 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 483 1483 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 514 75.5 41.2 35 53 484 1484 18866 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 514 75.5 41.2 35 53 485 1485 27365 27385 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.8 43 35 50.4 486 1486 18686 18715 51.7 50 3986 19217 19196 50.2 40.9 1.5 520 75.6 41.3 35 53.1 487 1487 3381 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.4 35 53.1 488 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.3 35 53.1 489 1489 3782 3801 51.3 50 3980 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35.3 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.3 50 3991 4444 4424 424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.2 525 0.9 44.6 35 53.1 475 1499 9999 9946 50 50 3992 13747 13726 50.8 40.9 0.7 663 75.5 40.6 35 53.1 475 1499 9999 9946 50 50 3994 10439 10431 50.9 474 0.9 521 75.4 40.9 35 52.9							3977	27568	27548	50.2	42.9	1.2	202	74.6	44.1	35	
489 1489 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 1.3 510 75.7 41.8 35 53.4 461 1481 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.3 213 75 44.6 35 52.9 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.4 463 1483 18704 18724 50.8 47.6 3983 19217 19196 50.2 40.9 0.5 512 75.5 41.2 35 53.4 464 1484 18696 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 514 75.5 41.2 35 53.4 465 1485 27365 27344 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.8 43 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.2 75.6 41.4 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.2 75.6 41.4 35 53.1 469 1489 3363 3361 3361 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.2 75.6 41.4 35 53.1 469 1489 3361 3361 3361 50.5 42.9 3987 3646 3625 52 40.9 1.5 52.2 75.6 41.4 35 53.1 470 1490 13039 3068 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 53.1 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 472 1492 3909 9946 50 50 3992 13757 13725 50.8 40.9 0.7 666 43.1 35 53.1 473 1493 2223 2243 50.2 42.9 3993 2747 2727 50 42.9 0.2 52.5 69.9 44.6 35 53.9 475 1499 9999 9946 50 50 39994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9								27571	27551	51.4	42.9	0	205	74.6		_	
461 481 27367 27385 51.4 52.6 3980 27576 27555 51 40.9 0.4 210 74.8 44.3 35 52.8 462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 51.2 75.5 41.2 35 53.3 463 1483 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 51.2 75.5 41.2 35 53.3 464 1484 18696 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 51.4 75.5 41.2 35 53.3 465 1485 27365 27384 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.5 64 33 50.3 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 526 75.6 41.3 35 53.1 468 1488 3381 3381 50.5 42.9 3988 3647 3628 50.6 45.0 1.5 52.6 75.5 43.7 35 53.1 469 1489 3782 3301 51.3 50 3980 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35.3 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52.1 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.7 60.2 57.6 44.3 35 53.1 475 1494 9929 9946 50 50 3994 10439 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9								9376	9355	51	40.9	1.3	510	75.7	41.8	35	
462 1482 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.9 463 1483 18704 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53.9 464 1484 18866 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 51.4 75.5 41.2 35 53.1 465 1485 27365 27384 52.6 50 3985 27464 27443 54.4 40.5 41.4 100 70.8 43 35 50.4 466 1486 18666 18715 51.7 50 3986 19215 19194 50.2 40.9 1.5 520 75.6 41.3 35 53.1 467 1487 3381 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.4 35 53.1 468 1488 3381 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 528 75.5 43.7 35 53.1 469 1489 3782 3301 51.3 50 3989 3647 3628 50.6 45 0.1 287 75.6 43.9 35 53.1 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.2 526 0.9 44.0 35 53.9 475 1499 9999 9946 50 50 3994 10439 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9									27555	51	40.9	0.4	210	74.8	44.3	35	
462 482 87/4 18724 50.8 47.6 3982 19215 19194 50.2 40.9 0.5 512 75.5 41.2 35 53 464 1484 18696 18715 51.7 50 3984 19215 19194 50.2 40.9 0.5 514 75.5 41.2 35 53 465 1485 27365 27364 52.6 50 3985 27464 27443 54 45.5 1.4 100 70.8 43 35 53.1 465 1486 1486 18696 18715 51.7 50 3986 19217 19196 50.2 40.9 1.5 520 75.6 41.3 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 522 75.6 41.4 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 528 75.5 41.4 35 53.1 469 1489 3393 3782 3801 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 470 1490 13039 13068 51.8 50 3989 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.2 525 69 44.5 53.3 474 1494 9929 9946 50 50 39394 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 475 1499 1499 1490 1400								27579	27558	51.1	40.9	0.3	213	75	44.6	35	
463 [1483] 18704 [18724] 50.8 [47.6] 9893 [19217] 19196 [50.2] 40.9 [0.5] 514 [75.5] 41.2 [35] 53 [465] 1485 [18736] 27364 [52.6] 50.3984 [19215] 19194 [50.2] 40.9 [1.5] 520 [75.6] 41.3 [35] 53.1 [465] 1485 [27365] 27364 [52.6] 50.3985 [27464] 27443 [54] 45.5 [1.4] 100 [70.8] 43 [35] 50.4 [466] 1486 [18696] 18715 [51.7] 50 [3986] 19217 [19196] 50.2 [40.9] 1.5 [522] 75.6 [41.4] 35 [53.1] 467 [487] 3361 [3381] 50.5 [42.9] 9387 [3646] 3625 [52] 40.9 [1.5] 522 [75.6] 41.4 [35] 53.1 [487] 488 [1488] 3361 [3331] 50.5 [42.9] 9388 [3647] 3628 [50.6] 45.0 [4.9] 40.9 [1.5] 522 [75.6] 41.4 [35] 53.1 [487] 488 [1488] 3361 [3331] 50.5 [42.9] 9388 [3647] 3628 [50.6] 45.0 [4.9] 0.7 [64] 75.5 [40.5] 35 [53.1] 469 [1489] 3782 [3801] 51.3 [50] 9399 [4445] 4425 [50.6] 42.9 [0.7] 647 [75.5] 40.5 [35] 53.1 [470] [490] 13039 [13058] 51.8 [50] 9399 [4444] 4424 [50.6] 42.9 [0.7] 663 [75.5] 40.6 [35] 52.1 [471] [491] 3782 [3801] 51.3 [50] 9391 [4444] 4424 [50.6] 42.9 [0.7] 663 [75.5] 40.6 [35] 53.1 [472] [492] [13040] [13059] 50.9 [50] 9392 [13747] [13766] 50.8 [40.9] 0.1 [70] 70.6 [43.1] 35 [53.1] 473 [493] 2223 [2243] 50.2 [42.9] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [1494] 9929 [9946] 50.5 [50] 9394 [10449] [10431] 50.9 [47.4] 0.9 [521] 75.4 [40.9] 35 [52.9] 475 [40.9] 90.9 [40.9] 40.9 [40.48] [40.48] [40.48] [40.48] [40.9] 90.9 [521] 75.4 [40.9] 35 [52.9] 475 [40.9] 90.9 [40.9] 40.9 [40.48] [40.48] [40.48] [40.9] 90.9 [40.9]								19215	19194	50.2	40.9	0.5	512	75.5	41.2	$\overline{}$	
465 1486 1896 18715 51.7 50 3984 19215 19194 50.2 40.9 1.5 520 75.6 41.3 35 53.1 465 1485 27385 27384 52.6 50 3985 27464 27443 27443 54 45.5 1.4 100 70.8 43 35 50.4 466 1486 18686 18715 51.7 50 3986 19217 19196 50.2 40.9 1.5 522 75.6 41.4 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 528 75.5 43.7 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 286 75.5 43.7 35 53.1 469 1489 3782 3801 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.2 526 0.3 44.9 34.3 53.3 474 1494 9929 9946 50 50 3994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 475 1497 1807 1									19196	50.2	40.9	0.5	514				
465 [1485 27365 27344 52.6 50]9965 27464 27443 54 45.5 1.4 100 70.8 43 35 50.4 466 [1486 18696 18696 18715 51.7 50 3966 19217 19196 50.2 40.9 1.5 522 75.6 41.4 35 53.1 467 [1487 3361 3381 50.5 42.9]987 3646 3625 52 40.9 1.5 286 75.5 43.7 35 53.1 468 [1488 3361 3381 50.5 42.9]988 3647 3628 50.6 45 0.1 287 75.6 43.9 35 53.1 470 [1490 19308 3762 3801 51.3 50]989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 471 [1491 3782 3801 51.3 50]989 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 472 [1492 13040 13058 50.8 50.9 50]989 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 473 [1493 2223 2243 50.2 42.9]993 2747 2727 50 42.9 0.2 525 60.9 44.0 35 53.9 474 [1494 9929 9946 50 50 39994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 475 [1495 18077 1807								19215	19194	50.2	40.9	1.5	520				
466 1486 16866 16715 51.7 50 3966 19217 19196 50.2 40.9 1.5 522 75.6 41.4 35 53.1 467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 286 75.5 43.7 35 53.1 468 1488 3361 3381 50.5 42.9 3987 3647 3628 50.6 45 0.1 287 75.6 43.9 35 53.1 469 1489 3782 3801 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 472 1492 13040 13059 50.3 50 3992 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3993 2747 2727 50 42.8 0.2 525 76.9 44.6 35 53.9 474 1494 9929 9946 50 50 3994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 475 1493 18077 18077 18077 18077 1878 10439 10439 10439 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9								27464	27443	54	45.5	1.4	100				
467 1487 3361 3381 50.5 42.9 3987 3646 3625 52 40.9 1.5 286 75.5 43.7 35 53.1 468 1489 3381 3381 50.5 42.9 3988 3647 3628 50.6 45 0.1 287 75.6 43.9 35 53.1 469 1489 3782 3801 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 473 1493 2223 2243 50.2 42.9 3983 2747 2727 50 42.9 0.2 525 76.9 44.6 35 53.9 474 1494 9929 9946 50 50 3994 10439 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 475 1494 9929 9946 50 50 3994 10439 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9								19217	19196	50.2	40.9	1.5					
468] 1488 3361 3381 50.5 42.9] 9988 3647 3628 50.6 45 0.1 287 75.6 43.9 35 53.1 469 1489 3782 3801 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 470 1490 13039 13058 51.8 50] 9990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 472 1492 13040 13059 50.9 50 3992 13747 13726 50.8 40.9 0.1 708 76.6 43.1 35 54 473 1493 2223 2243 50.2 42.9] 9993 2747 2727 50 42.9 0.2 525 60.9 44.0 35 53.9 474 1494 9929 9946 50 50 3994 10449 10431 50.9 474 0.9 521 75.4 40.9 35 52.9							3987	3646	3625	52	40.9	1.5	286				
469] 1489 3782 3801 51.3 50 3989 4445 4425 50.6 42.9 0.7 664 75.5 40.5 35 53.1 470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.8 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 472 1492 13040 13059 50.9 50 3992 13747 13726 50.8 40.9 0.1 708 76.6 43.1 35 54 473 1493 2223 2243 50.2 42.9 3993 2747 2727 50 42.9 0.2 525 76.9 44.6 35 53.9 474 1494 9929 9946 50 50 3994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9									3628	50.6	45	0.1	287				
470 1490 13039 13058 51.8 50 3990 13155 13137 52.1 52.6 0.3 117 73.4 47 35 52 471 1491 3782 3801 51.3 50 3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 472 1492 13040 13059 50.9 50 3992 13747 13726 50.8 40.9 0.1 708 76.6 43.1 35 53.9 474 1494 9929 9946 50 50 3994 10439 10431 50.9 47.4 0.9 52 76.9 44.6 35 53.9 475 1493							3989	4445	4425	50.6	42.9	0.7				_	
471 [1491 3782] 3801 51.3 50 [3991 4444 4424 50.6 42.9 0.7 663 75.5 40.6 35 53.1 472 [1492 13040 13059 50.9 50] 3992 13747 13726 50.8 40.9 0.1 708 76.6 43.1 35 53.1 473 [1493 2223 2243 50.2 42.9] 3993 2747 2727 50 42.9 0.2 525 6.9 44.6 35 53.9 474 [1494 9929] 9946 50 50 50 [3994 10439 1043] 50.9 474 0.9 521 75.4 40.9 35 52.9 475 [1495 18077] 18077 [1497 1516]						50	3990	13155	13137	52.1	52.6						
472 [1492 13040 13069 50.9 50 [3992 13747 13726 50.8 40.9 0.1 708 76.6 43.1 35 54 473 1493 2223 2243 50.2 42.9 [3993 2747 2727 50 42.9 0.2 52.5 76.9 44.6 35 53.9 474 494 9929 9946 50 50 [3994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 47.5 48.9 18977 48.9 48.7 48.								4444	4424	50.6	42.9	0.7	663				
473 1493 2223 2243 50.2 42.9 3993 2747 2727 50 42.9 0.2 525 76.9 44.6 35 53.9 474 1494 9929 9946 50 50 50 3994 10439 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9 47.5 1495 18977 18907 51.5 72.0000 47.7 14007 51.5 72.0000 47.7							3992	13747	13726	50.8	40.9	0.1					
474 1494 9929 9946 50 50 3994 10449 10431 50.9 47.4 0.9 521 75.4 40.9 35 52.9							3993	2747	2727	50	42.9						
4/5 1495 18077 19007 51 5 47 6 2005 40706 40706								10449	10431	50.9	47.4	0.9	521				
	475	1495	18077	18097	51.5	47.6	3995	18702	18685	50.2	50	1.4	626				

476	1496	3360	3379	50.7	4	3996	3646	3625	52	40.9	1.3	287	75.4	43.6	35	53.1
477	1497	26708	26731	54.2	41.	3997	27463	27443	52.7	42.9	1.5				35	
478	1498	26708	26731	54.2	41.	7 3998	27464	27444	53		1.2	757	75.9			
479	1499	26708	26731	54.2	41.	3999	27464	27445	52.4	45	1.8	757	75.9			
480	1500	4	22	52.3	52.6	4000	713	695		47.4	1.6	710				54 55.7
481	1501	26708	26727	50	48	4001	27462	27443		45	1.3	.755	75.9		35	53.2
482	1502	3360	3380	51.4	42.9	4002	3646	3625	52	40.9	0.6	287	75.4	43.6	_	
483	1503	26708	26727	50	45		27463	27445	50.8	42.1	0.8	756	75.9	43.0	35	53.3 53.2
484	1504	988	1006	52.2	52.6	4004	1493	1474	50.8	45	1.5	506	76.5	43.7		
485	1505	12352	12375	52.9	41.7		12993	12975	51.4	47.4	1.5	642	76.5		35	53.9
486	1506	3360	3380	51.4	42.9		3647	3628	50.6	45	0.8	288	75.5	43	35	. 54
487	1507	18074	18094	51.1	42.9		18702	18685	50.0	50	0.8	629		43.8	35	53.1
488	1508	3360	3380	51.4	42.9	1	3650	3631	53.1	50	1.7		76.2	42.3	35	53.5
489		8374	8395	52.4	45.5		8928	8911	51.9	50		291	75.6	44	35	53.5
490	1510	9929	9946	50	50		10449	10428	51.9	40.9	0.6 1.9	555 521	75.1	40	35	53.2
491	1511	26421	26441	51.5		4011	27132	27111	50.3	40.9	1.9		75.4	40.9	35	52.9
		10250	10274	51.6	40		10356	10336	52.4	47.6		712	77.1	44.2	35	54.2
	1513	18074	18093	50.3	45		18702	18685	50.2	47.0 50	0.8	107	70.8	42.1	35	50.2
		18017	18036	54.8	55		18220	18202	54.8	52.6		629	76.2	42.3	35	53.5
495	1515	18017	18036	54.8	55		18225	18206	53.7	52.6	0	204	74.3	43.1	35	53.5
		18017	18036	54.8		4016	18232	18210	54.4	47.8	1.1	209	74.3	43.1	35	53.2
497	1517	18017	18036	54.8	55		18234				0.4	216	74.6	43.5	35	53.6
	1518	18017	18036	54.8		4018	18235	18214 18215	53.4 54.2	47.6	1.4	218	74.7	43.6	35	53.4
_	1519	18017	18036	54.8	55		18443			52.4	0.6	219	.74.8	43.8	35	53.7
	1520	18012	18031	53.2	55		18220	18424	55.9	55	1.1	427	76	43.1	35	54.7
	1521	18013	18031	50.6	52.6		18223	18202	54.8	52.6	1.7	209	74.5	43.5	35	53.2
	1522	18013	18031	50.6	52.6		18231	18206	51.8	50	1.1	211	74.4	43.1	35	52.4
	1523	18013	18031	50.6	52.6		18233	18210	52.2	45.5	1.6	219	74.6	43.4	35	52.5
	1524	18013	18031	50.6	52.6		18233	18214	52	50	1.4	221	74.8	43.9	35	52.7
	1525	18013	18031	50.6		4024		18215	51.3	52.6	0.7	221	74.8	43.9	35	52.7
	1526	18009	18029	53.3	52.4	4025	18662 18220	18641	50.4	40.9	0.2	650	76.3	42.6	35	53.7
	1527	18011	18029	51.3	52.4			18202	54.8	52.6	1.6	212	74.5	43.4	35	53.2
	1528	18011	18029	51.3	52.6	4027	18223	18206	51.8	50	0.5	213	74.4	43.2	35	52.6
	1529	18011	18029	51.3		4028	18231 18233	18210	52.2	45.5	0.9	221	74.7	43.4	35	52.8
	1530	18011	18029	51.3		4029		18214	52	50	0.7	223	74.9	43.9	35	52.9
	1531	16374	16397	52.8		4030	18233	18215	51.3	52.6	0	.223	74.9	43.9	35	52.9
	1532	16378	16397	50.4		4031		16751	53.6	41.7	0.8	401	. 75	40.9	35	53.4
	1533	2223	2243	50.4		4032	16780 2997	16760	51.4	42.9	1	403	75	40.9	35	52.7
	1534	2428	2447	51.5				2976	51.4	40.9	1.2	775	76.7	43.1	35	53.9
	1535	16548	16566	54.9		4034 4035	3082	3058	52.3	40	0.8	655	76.3	42.6	35	54
	1536	16367	16386	51.4		4035	16774 16774	16751	53.6	41.7	1.3	227	73.9	41.4	35	52.9
	1537	3230	3249	50.1		4036		16752	52.2	43.5	0.8	408	75	40.9	35	53
	1538	8221	8240	52.4			3497	3478	51.3	50	1.2	268	74.4	41.4	35	52.2
	1539	3232	3252	51.1		4038	8920	8901	53.4	50	1	700	75.3	40	35	53.6
	1540	3232	3252			4039	3500	3481	51.2	50	0.1	269	74.5	41.6	35	52.5
	1540	16367		51.1		4040	3497	3478	51.3	50	0.2	266	74.5	41.7	35	52.6
			16386	51.4		4041	17111	17090	51.1	40.9	0.3	745	76.3	42.1	35	53.8
	1542 1543	16366 9930	16385	52.9		4042	16774	16751	53.6	41.7	0.8	409	75.1	41.1	35	53.5
			9948	51.5		4043	10670	10649	51.3	40.9	0.2	741	75.8	40.9	35	53.5
	1544 1545	12370 25354	12388	50.1		4044	12996	12977	50.2	40	0.2	627	76.4	42.7	35	53.6
			25372	50.9		4045	25650	25631	51.3	45	0.4	297	74.5	41.1	35	52.5
526 1	040	25354	25372	50.9	52.6	4046	25651	25634	50.4	50	0.5	298	74.6	41.3	35	52.4

	-	07 4 = 4															
		27 1547					6 4047	2577	2 2575	3 51.	9 50)	1 41	9 74.	8 40.	3 3	5 52.8
		28 1548					9 4048	150	1 148	2 50.	5 4					6 3	
		29 1549			2 50.	9 52.	6 4049	2583	1 2580								
		30 1550			5 50.	9 47.	4 4050	443	4 4410						-		
	5		2535	2537	2 50.	9 52.	6 4051	2583									
	53	32 1552	379	7 381	5 50.	9 47.	4 4052	443			_			_			
	53	33 1553	2448	1 2450	0 50.	1 4	5 4053	2493				-					
	53	34 1554	2534	8 2536	6 51.		4 4054	2583								_	
	53	35 1555	2534	8 2536	6 51.		4 4055	2583									
	53	36 1556	2441	9 2444	0 52.		4056	25080									
	53	7 1557	2442	0 2444				24527									
	53	8 1558	2534				0 405B	25650				0.3		1		_	
	53	9 1559	2534					25651		51.3		0.9	30:				52.4
- 1	54		2534					25831		50.4		0.1	304			1 35	52.5
	54	1 1561	2534									1	484	1		35	52.7
1	54		28618				1.00.	25831	25810	50.7	45.5	0.3	484		40.3	35	52.7
Ì	54		8867					29298		51.4		1.1	681		47.3	35	55.3
t	54		28820					9317	9297	50.5	42.9	1.8	451		41.7	35	53.1
t	54		27365				1	29301	29282	55.3	55	1.6	482		45.4	35	55.2
ŀ	54		28820					27464	27443	54	45.5	0.8	100	70.8	43	35	50.6
t	54		28820					29306	29288	53.5	52.6	0.8	487	77.1	45.4	35	55.1
ŀ	54		28821	28840				29301	29282	55.3	55	1	482	77.1	45.4	35	55.4
ŀ	549		27370					29306	29288	53.5	52.6	1.7	486	77.1	45.3	35	54.6
t		0 1570	28820		54.8		4069	27675	27656	50	· 40	0.1	306	74.2	40.2	35	52
H	55		27370		50.1	47.6		29301	29282	55.3	55	0.4	482	77.1	45.4	. 35	55.5
H	552		2429	2447	50.1	45	4071	27674	27654	51.9	42.9	1.8	305	74.2	40.3	35	52.1
H		1573	27375			47.4		3188	3167	50.2	40.9	0	760	76.6	42.9	35	53.8
H	554		27375	27392	50	50	4073	27675	27656	50	40	0	301	74.1	40.2	35	52
H	555		19795		50	50	4074	27674	27654	51.9	42.9	1.9	300	74.2	40.3	35	52
H		1576	3168	19814	50.4	45	4075	19916	19895	50.2	40.9	0.2	122	71.8	42.6	35	50.5
H	557		3168	3189	51		4076	3646	3625	52	40.9	1.1	479	75.7	42	35	53.4
\vdash		1578	18011	3189	51	45.5		3647	3628	50.6	45	0.4	480	75.8	42.1	35	53.3
H	559			18029	51.3	52.6		18662	18641	50.4	40.9	0.9	652	76.3	42.6	35	53.7
H	560		985	1004	51.1		4079	1493	1474	50.8	45	0.3	509	76.5	43.6	35	53.9
H	561	1581	12965	12985	51.2	42.9		13547	13528	50.2	45	0.9	583	77.3	45.3	35	54.3
H	562		2427	2445	52.1		4081	3188	3167	50.2	40.9	1.9	762	76.7	43	35	53.8
H	563	1582 1583	3360	3381	52.1		4082	3650	3631	53.1	50	1	291	75.6	44	35	53.7
H			12726	12746	51.3	47.6		12911	12892	50.5	50	0.8	186	73.5	41.9	35	51.7
H	564 565	1584	19800	19817	50.4		4084	19917	19896	50.9	45.5	0.5	118	71.9	43.2	35	50.6
H	566	1585	1402	1426	54.1		4085	1501	1478	54.6	41.7	0.5	100	72	46	35	51.8
H			2427	2445	52.1		1086	3082	3058	52.3	40	0.2	656	76.4	42.7	35	54.2
L	567	1587	8867	8887	52.3	47.6	1087	9257	9238	50.5	45	1.8	391	75.1	41.2	35	52.8
-	568	1588	8867	8887	52.3	47.6	1088	9249	9231	50.8	47.4	1.5	383	75.2	41.5	35	53
L		1589	8374	8394	51	42.9	1089	8928	8911	51.9	50	0.8	555	75.1	40	35	53
L		1590	8867	8887	52.3	47.6	1090	9249	9230	51.5	45	0.8	383	75.2	41.5	35	53.2
L		1591	28964	28984	54.3	52.4	091	29301	29282	55.3	55	11	338	76.5	45.3	35	
L		1592	8867	8887	52.3	47.6	092	9249	9229	53	47.6	0.6	383	75.2	41.5		54.9
L		1593	12962	12980	50.7	47.4	093	13547	13528	50.2	45	0.5	586	77.4		35	53.4
L		1594	9931	9950	50.2	45 4	094	10605	10588	51.1	50	0.9	675	75.8	45.4 41.2	35	54.3
L		1595	19801	19819	53.2	52.6 4	095	19918	19896	52.2	43.5	1	118	71.9	43.2	35	53.2
L		1596	9055	9079	52.8	40 4		9376	9355	51	40.9	1.8	322	75.1		35	51.1
L	577	1597	19878	19899	50.5	40.9 4	097	20033	20016	50.4	50	0.1	156	73.4	42.2	35	53
										-0.4	- 50	J.11	130	13.4	43.6	35	51.6

578 1598 17608 17628 50.9 42.9 4068 18233 18214 52 50 1.1 626 75.3 40.3 35 53.2 589 1690 29179 29199 51.4 42.9 4100 29358 29339 52.8 50 1.4 180 74.8 45.6 35 52.9 581 1601 29182 29202 51.2 42.9 4101 29358 29339 52.8 50 1.4 180 74.8 45.6 35 52.9 582 1602 4 22 52.3 52.6 4102 29558 29339 52.8 50 1.4 180 74.8 45.6 35 52.9 583 1603 8221 8240 52.4 50 4103 8920 8902 52.8 52.6 3.7 70.0 72.9 40.4 35 54.6 583 1603 8221 8240 52.4 50 4104 16774 16751 53.6 41.7 0.1 22.1 73.7 41.2 55.5 585 1605 16554 5657 53.7 52.6 4106 16774 16761 16751 53.6 41.7 0.1 22.1 73.7 41.2 55.6 586 1605 16555 16572 53.7 52.6 4106 29412 29393 50.3 45.5 0.3 227 75.4 44.9 35 53.6 586 1605 29168 29205 50.1 40 4106 29412 29393 50.3 45.5 0.3 227 75.4 44.9 35 53.6 588 1608 29182 29205 54.6 41.7 4108 29358 29339 52.8 50 1.7 177 74.6 45.2 35 53.6 589 1609 4 22 22.3 52.6 4109 255 235 51.3 47.6 1.1 227 73.4 41.3 55.2 590 1610 3230 3249 50.1 45 4110 3950 3481 51.2 50 1.1 271 74.4 41.3 35 52.2 590 1611 13040 13059 50.9 50.4 50.4 111 3177 13156 50.6 47.4 40.9 50.5 138 73.7 45.7 36.5 590 1613 19995 20012 50.4 50.4 111 20151 20597 50.6 47.4 10.9 50.7 75.8 40.1 35 53.5 590 1616 12370 12388 50.1 47.4 4115 20151 20597 50.6 47.4 1.2 20.1 75.3 40.1 35 53.5 590 1616 12370 12388 50.1 47.4 4115 20152 20597 50.6 47.4 1.2 20.1 75.3 40.1 35 53.5 590 1616 12370 12388 50.1 47.4 4115 20152 20597 50.6 47.4 1.2 60.0 75.8 40.1 35 53.5 590 1616 12370 12388 50.1 47.4 4115																
Sep 1600	578	1598	17608	17628	50.9	42.9 4098	18233	18214	52	50	1.1	626	75.3	40.3	35	53.1
561 1601 29182 29202 51.2 42.9 I4101 29368 29339 52.8 60 1.6 177 74.6 45.2 35 52.7 582 1602 4 22 52.3 52.6 I4102 253 233 51.8 47.6 0.5 250 37.0 75.2 40.4 33 53.6 584 1604 16564 16572 53.3 50.4103 8620 869.2 52.8 52.6 0.3 700 75.3 40 35 55.6 585 1605 1605 16572 53.3 50.4110 16774 16751 53.6 41.7 0.1 221 75.7 74.4 41.2 33.3 53.6 41.7 0.1 221 25.2 50.0 1.7 177 74.6 42.2 53.3 33.3 53.6 0.1 62.4 75.4 44.2 35.5 55.2 55.0 1.7 177 74.6 42.6 23.5 55.2 55.2 55.2	579	1599	17608	17627	50.2	45 4099	18233	18214	52	50	1.8	626	75.3	40.3	35	52.9
S82 1802	580	1600	29179	29199	51.4	42.9 4100	29358	29339	52.8	50	1.4	180	74.8	45.6	35	52.9
S83 1603 8821 8240 52.4 50.4 4103 8920 8902 82.8 52.6 0.3 700 75.3 40 35 53.6 584 1604 16654 16572 53.7 52.6 4104 16774 16691 51 42.9 0.7 157 73.4 41.2 35 52.8 585 1606 6565 16572 53.3 50.4 40.4 106 29412 29393 50.3 45 0.3 227 75.4 44.9 35 53.6 586 1606 29186 29205 50.1 40.4 4106 29412 29393 50.3 45 0.3 227 75.4 44.9 35 53.6 587 1607 2429 2447 50.2 47.4 4107 3052 3033 50.3 50 0.1 624 76.3 42.6 35 53.6 588 1608 29182 29205 54.6 41.7 4108 29368 29339 52.8 50 1.7 177 74.6 45.2 35 53.6 589 1609 4 22 52.3 52.6 4109 255 225 51.3 47.6 1 252 76.3 46.4 52 35 53.6 590 1610 3230 3249 50.1 45.4 4110 3050 3481 15.2 50 1.1 271 74.4 41.3 35 52.9 591 1611 3040 13059 50.9 50.4 4112 17039 17022 51.4 50 0.3 489 75.8 42.1 35 53.5 591 1612 16551 16568 51.1 50.4 112 17039 17022 51.4 50 0.3 489 75.8 42.1 35 53.5 591 1614 19995 20013 51.8 52.6 4114 20615 20597 50.6 47.4 12 621 75.3 40.1 35 53.5 591 1616 12370 12386 50.1 47.4 4114 20615 20597 50.6 47.4 12 621 75.3 40.1 35 53.6 591 1616 8374 8393 51.2 45 4116 8928 8911 51.9 50 0.7 555 75.1 40 35 53.6 591 1617 24174 24199 50.9 42.9 4117 24936 24919 51.8 50 0.7 758 75.8 41 35 53.5 599 1619 7679 7696 50.6 50.4 111 8049 8032 50.4 40.9 0.7 758 75.8 41 35 53.5 599 1619 7679 7696 50.6 50.4 111 24936 24919 51.8 50 0.7 758 75.8 41 35 53.5 599 1619 7679 7696 50.6 50.4 119 8049 8032 50.4 50 0.7 758 75.8 41 35 53.5 599 1619 7679 7696 50.6 50.4 119 8049 8032 50.4 50 0.7 758 75.8 41 35 53.5 50.	581	1601	29182	29202	51.2	42.9 4101	29358	29339	52.8	50	1.6	177	74.6	45.2	35	52.7
Sept 1004 16564 16572 53.7 52.6 4104 16774 16751 53.6 41.7 0.1 221 73.7 41.2 35 52.6 5651 16055 16552 50.3 50 4105 16711 168691 51 42.9 0.7 175 73.4 43.3 35 51.6 568 1606 29186 29050 50.1 40 4106 29412 29383 0.3 45 0.3 227 75.4 44.9 35 52.9 597 1607 2429 2447 50.2 47.4 4108 29362 29303 50.3 50 0.1 624 76.3 42.6 35 53.6 588 1608 29182 29205 54.6 41.7 4108 29368 29338 52.8 50 1.7 77.7 74.6 45.2 35 53.6 588 1608 29182 29205 54.6 41.7 4108 29368 29338 52.8 50 1.7 77.7 74.6 45.2 35 53.6 588 1608 29182 29205 54.6 41.7 4108 29368 29338 52.8 50 1.7 177 74.6 45.2 35 53.5 598 1610 3320 3249 50.1 45.4110 3500 3481 51.2 50 1.1 271 74.4 41.3 35 52.2 591 1611 13040 13059 50.9 50.4111 13177 13156 50.4 40.9 0.5 138 73.7 45.7 35 51.8 593 1613 19995 20012 50.4 50.4111 20615 20597 50.6 47.4 0.2 621 75.3 40.1 35 52.9 594 1614 19996 20013 51.8 52.4 1114 24938 24919 51.8 50 0.3 449 75.8 42.1 35 53.5 598 1615 12370 12388 50.1 47.4 4115 12993 12975 51.4 47.4 1.3 624 76.4 42.9 35 53.6 599 1619 24179 24198 51 45.4118 24936 24919 51.8 50 0.7 75.8 75.8 41.3 53.5 599 1619 24179 24198 51 45.54118 24936 24919 51.8 50 0.7 75.8 75.8 41.3 53.5 53.6 50.1 12370 7679 7698 50.6 50.4113 24938 24919 51.8 50 0.7 75.8 75.8 41.3 53.5 599 1619 24179 24200 53.3 40.9 4122 24938 24919 51.8 50 0.7 75.8 75.8 41.3 53.5 599 1619 24179 24206 53.3 40.9 4122 24938 24919 51.8 50 0.7 75.8 75.8 41.3 53.5 50.0 1620 13177 13197 50.3 60.1412 24938 24919 51.8 50 0.7 75.8 75.8 41.3 53.5 53.6 6	582	1602	4	22	52.3	52.6 4102	253	233	51.8	47.6	0.5	250	76.2	46.4	35	54
Self 1605 16556 16572 50.3 50 4105 16711 16669 51 42.9 0.7 157 73.4 43.3 35 51.6	583	1603	8221	8240	52.4	50 4103	8920	8902	52.8	52.6	0.3	700	75.3	40	35	53.6
See 1606 29186 29205 50.1 40 4106 29412 29393 50.3 45 50.3 227 75.4 44.9 35 52.9 567 1607 2429 2447 50.2 47.4 4107 3052 3033 50.3 50 0.1 624 76.3 42.6 35 53.6 588 1608 29182 29205 54.6 41.7 4108 29385 29393 52.8 50 0.1 717 77 46 45.2 35 53.6 589 1609 4 22 52.3 52.6 4109 255 236 51.3 47.6 11 252 76.3 46.4 35 53.9 590 1610 3230 3245 50.1 45 4110 3500 3481 51.2 50 1.1 271 74.4 41.3 35 52.2 591 1611 13040 13059 50.9 504111 13177 13166 50.4 40.9 0.5 138 73.7 45.7 35 52.5 592 1612 16551 16568 51.1 50 4112 17099 17022 51.4 50 0.3 489 75.8 42.1 35 53.5 593 1613 19995 20013 51.8 52.6 4114 20615 20597 50.6 47.4 0.2 621 75.3 40.1 35 52.9 594 1614 19985 20013 51.8 52.6 4114 20615 20597 50.6 47.4 1.2 621 75.3 40.1 35 52.9 595 1615 12370 12388 50.1 47.4 4115 12993 12975 51.4 47.4 1.3 624 76.4 42.9 35 53.6 599 1617 24174 24194 50.9 42.9 4117 24936 24919 51.8 50 0.7 755 75.8 41 35 53.5 599 1619 7679 7698 50.6 50 4119 80.98 8911 51.9 50 0.7 756 75.8 41 35 53.5 599 1619 7679 7698 50.6 50 4119 80.99 80.8 50.4 50.9 10.2 37.7 75.8 41 35 53.5 600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 144 73.2 43.8 35 53 600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 144 73.2 43.8 35 53 600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 144 73.2 43.8 35 53 600 1620 13177 13197 50.3 42.9 4120 42420 53.3 50.9 50.9 50.9 50.9 75.7 40.9 35 53 50.0 40.9 50.9 50.9 50.9 50.9 50.9 50.9 50.9 50.9 50.9 50.9 50.9 50.9 50	584	1604	16554	16572	53.7	52.6 4104	16774	16751	53.6	41.7	0.1	221	73.7	41.2	35	52.8
586 1606 29186 29205 50.1 40.4106 29412 29393 50.3 45 0.3 227 75.4 44.9 35 52.9 587 1607 2429 2447 50.2 47.4 4107 3052 3033 50.3 50 0.1 624 76.3 42.6 35 53.6 588 1808 29182 29205 54.8 41.7 4108 29358 29399 52.8 50 1.7 177 74.6 45.2 35 53.2 599 1609 4 22 52.3 52.6 4109 255 235 51.3 47.6 1 252 76.3 46.4 35 53.9 590 1610 3220 3249 50.1 45 4110 3500 3481 51.2 50 1.1 271 74.4 41.3 35 52.2 591 1611 13040 13059 50.9 50 4111 13177 13186 50.4 40.9 0.5 138 73.7 45.7 35 51.8 592 1612 16551 16568 51.1 50 4112 17039 17022 51.4 50 0.3 489 75.8 40.1 35 52.5 593 1613 13995 20012 50.4 50 4113 20615 20597 50.6 47.4 0.2 621 75.3 40.1 35 53.5 595 1615 12370 12388 50.1 47.4 4115 12993 12975 51.4 47.4 1.3 624 76.4 42.9 36 53.6 599 1617 24174 24194 50.9 42.9 4117 24936 24919 51.8 50 0.7 555 75.8 41 35 53.5 599 1617 24174 24194 50.9 42.9 4118 24936 24919 51.8 50 0.6 763 75.8 41 35 53.5 600 1620 13177 13197 50.3 42.9 4118 24936 24919 51.8 50 0.7 755 75.8 41 35 53.5 600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 14.4 73.2 43.8 35 53.6 601 1621 24170 24200 53.3 40.9 4121 24934 24913 53.4 45.5 0.2 756 75.8 411 35 53.5 603 1623 2427 2445 52.1 52.6 4123 3052 3033 50.3 50.1 47.6 1.1 47.7 42.9 43.8 35 53.6 603 1623 2427 24435 50.4 52.6 4123 3052 3033 50.3 50.1 47.6 1.1 1.1 1.1 1.1 1.3 53.2 603 1628 18074 18094 51.1 42.9 4128 1862 18611 50.2 41.1 10.0 71.3 42.2 36 53.6 604 1624 24418 24438 50.4 50.4 4116 8998 8	585	1605	16555	16572	50.3	50 4105	16711	16691	51	42.9	0.7	157	73.4	43.3	35	51.6
588 1608 29182 29205 54.6 41.7 4108 29388 29339 52.8 50 1.7 177 74.6 48.2 35 52.3 52.8 6109 25 525 51.3 47.6 1 252 76.3 46.4 35 53.9 590 1610 3220 3249 50.1 484110 3500 3481 51.2 50 1.1 277.7 46.7 435 55.2 591 1611 13040 13059 50.9 50 4111 13177 13156 50.4 40.9 0.5 138 73.7 45.7 35 51.8 592 1612 16551 16668 51.1 504112 17039 17022 51.4 40.9 0.5 138 73.7 45.7 35 51.8 593 1613 13995 20012 50.4 1411 17177 13156 50.4 47.4 1.2 621 75.3 40.1 35 53.3 <	586	1606	29186	29205	50.1	40 4106	29412	29393	50.3	45	0.3	227	75.4	44.9	35	
S89 1609 4 22 52.3 52.6 4109 255 235 51.3 47.6 1 252 76.3 46.4 35 53.9	587	1607	2429	2447	50.2	47.4 4107	3052	3033	50.3	50	0.1	624	76.3	42.6	35	53.6
Sep 1610 3230 3249 50.1 45 4110 3500 3481 51.2 50 1.1 271 74.4 41.3 35 52.2 591 1611 13040 13059 50.9 50.4 111 13177 13156 50.4 40.9 0.5 138 73.7 45.7 35 51.8 502 1612 16551 16568 51.1 50 4112 17039 17022 51.4 50 0.3 489 75.8 42.1 35 53.5 53	588	1608	29182	29205	54.6	41.7 4108	29358	29339	52.8	50	1.7	177	74.6	45.2	35	53.2
599 1610 3230 3249 50.1 45 4110 3500 3481 51.2 50 1.1 271 74.4 41.3 35 52.2	589	1609	4	22	52.3	52.6 4109	255	235	51.3	47.6	1	252	76.3	46.4	35	53.9
Sep 1612 16551 16568 51.1 50 4112 17039 17022 51.4 50 0.3 489 75.8 42.1 35 53.5	590	1610	3230	3249	50.1	45 4110	3500	3481	51.2	50	1.1	271	74.4	41.3	35	
S83 1613 19995 20012 50.4 50 4113 20615 20597 50.6 47.4 0.2 621 75.3 40.1 35 52.5	591	1611	13040	13059	50.9	50 4111	13177	13156	50.4	40.9	0.5	138	73.7	45.7	35	51.8
596 1614 19995 20013 51.8 52.6 4114 20615 20597 50.6 47.4 1.2 621 75.3 40.1 35 535 585 1615 12370 12388 50.1 47.4 4115 12993 12975 51.4 47.4 1.3 624 76.4 42.9 36 55.8 596 1616 8374 8393 51.2 65 4116 8928 8911 51.9 50 0.7 555 75.1 40.3 55 589 1617 24174 24194 50.9 42.9 4117 24936 24919 51.8 50 0.8 763 75.8 41 35 53.5 599 1617 7617 7698 50.6 50 4119 8049 8032 50.4 50 0.2 371 75.4 42.3 35 53 600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 144 73.2 43.8 35 51.4 601 1621 24176 24200 53.3 40.9 4121 24934 24919 51.8 50 0.7 758 75.8 41 35 53.5 600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 144 73.2 43.8 35 51.4 601 1621 24176 24200 53.3 40.9 4121 24934 24919 53.4 45.5 0.2 756 75.8 41 35 54.5 602 1622 9927 99945 50.8 52.6 4122 10670 10649 51.3 40.9 0.5 74.6 75.7 40.9 35 53.4 603 1623 2427 2445 52.1 52.6 4123 3052 3033 50.3 50 1.8 626 76.4 42.8 35 53.6 604 1624 24418 24438 50 47.4 4124 24527 24507 24494 53.2 41.7 0.6 101 71.1 43.6 53.5 607 1627 24418 24438 52.6 504 125 24517 24494 53.2 41.7 0.6 101 71.1 43.6 53.5 50.6 608 1626 8375 8396 51.8 45.5 4126 8929 8911 53.4 52.6 1.6 555 75.1 40 35 53.6 609 1629 18074 18094 51.1 42.9 4129 18632 18611 50.2 40.9 0.9 559 76.2 42.8 36 53.5 600 1623 18074 18094 51.1 42.9 4129 18632 18611 50.2 40.9 0.9 559 76.2 42.8 36 53.5 610 1630 13231 13251 50.1 42.9 4130 13545 13526 50.3 52.6 0.6 633 75.8 41.2 35 54.6 611 633 3792 3811 54 55 4132 4446 4424 52.4 43.5 0.6 655 75	592	1612	16551	16568	. 51.1	50 4112	17039	17022	51.4	50	0.3	489	75.8	42.1	35	53.5
594 1614 19985 20013 51.8 52.6 4114 20615 20597 50.6 47.4 1.2 621 75.3 40.1 35 53 595 1615 12370 12388 50.1 47.4 4115 12993 12975 151.4 47.4 13 624 76.4 42.9 35 50 50 1617 24174 24194 50.9 42.9 4117 24936 24919 51.8 50 0.8 763 75.8 41 35 53.5 599 1617 24179 24198 51.4 420 13320 151.8 50 0.7 788 75.8 41 35 53.5 509 1619 7679 7698 50.6 50 4119 8049 8032 50.4 50 0.7 788 75.8 41 35 53.5 600 1620 12177 13197 50.3 42.9 4120 13320 131.4 47.6 1.1	593	1613	19995	20012	50.4	50 4113	20615	20597	50.6	47.4	0.2	621	75.3	40.1	35	52.9
Sep 1616 8374 8393 51.2 45 4116 8928 8911 51.9 50 0.7 555 75.1 40 35 53	594	1614	19995	20013	51.8	52.6 4114	20615	20597	50.6	47.4	1.2	621	75.3	40.1	35	
596 1616	595	1615	12370	12388	50.1	47.4 4115	12993	12975	51.4	47.4	1.3	624	76.4	42.9	35	53.6
S98 1618 24179 24198 51 45 4118 24936 24919 51.8 50 0.7 758 75.8 41 35 53.5	596	1616	8374	8393	51.2	45 4116	8928	8911	51.9	50	0.7	555	75.1	· 40	35	
Sep 1619 7679 7698 50.6 50 4119 8049 8032 50.4 50 0.2 371 75.4 42.3 35 53.4	597	1617	24174	24194	50.9	42.9 4117	24936	24919	51.8	50	0.8	763	75.8	41	35	53.5
600 1620 13177 13197 50.3 42.9 4120 13320 13300 51.4 47.6 1.1 144 73.2 43.8 35 51.4 601 1621 24176 24200 53.3 40.9 4121 24934 24913 53.4 45.5 0.2 756 75.8 41 35 54.2 602 1622 9927 9945 50.8 52.6 4122 10670 10469 51.3 40.9 0.5 74.6 75.7 40.9 35 53.4 603 1623 2427 2445 52.1 52.6 4123 3052 3033 50.3 50 1.8 626 76.4 42.8 35 53.6 604 1624 24418 24436 52.6 50 4125 24577 24494 53.2 41.7 0.6 101 71.1 43.6 52.6 605 1625 24417 24436 52.6 50 4125 24517 24494 53.2 41.7 0.6 101 71.1 43.6 55 50.6 606 1626 8375 8396 51.8 45.5 4126 8929 8911 53.4 52.6 1.6 555 75.1 40 35 53.2 607 1627 24418 24439 52.9 45.5 4127 25080 25062 53.5 52.6 0.6 663 75.8 41.2 35 54 609 1628 18074 18094 51.1 42.9 4129 18632 18611 50.2 40.9 0.9 559 76.2 42.8 36 53.5 610 1630 13231 13251 50.1 42.9 4129 18632 18611 50.2 40.9 0.9 559 76.2 42.8 36 53.5 612 1632 3792 3811 54 55 4132 4446 4424 52.4 43.5 1.6 665 75.5 40.6 36 33.6 613 1633 25782 25805 52.1 41.7 4133 26182 26161 51.2 40.9 0.9 401 74.7 40.1 36 52.7 614 1634 13320 13251 52.4 45.5 4132 4446 4424 52.4 43.5 1.6 665 75.5 40.6 36 53.6 619 1639 25782 25805 52.1 41.7 4133 26182 26161 51.2 40.9 0.9 401 74.7 40.1 36 52.7 614 1634 13320 13251 52.4 45.5 4132 4446 4424 52.4 43.5 1.6 665 75.5 40.6 36 53.6 619 1639 1376 13196 51.1 50 4135 1480 1462 51.6 47.4 0.5 366 76.4 42.2 36 53.5 619 1639 1376 13196 51.4 47.6 4139 13545 13526 52.9 55 0.2 316 77 47.1 47.2 36 54.8 619 1639 1376 13196 51.4 47.8 413	598	1618	24179	24198	51	45 4118	24936	24919	51.8	50	0.7	758	75.8	41	35	53.5
601 1621 24179 24200 53.3 40.9 4121 249.34 24913 53.4 45.5 0.2 756 75.8 41 35 54.2 602 1622 9927 9945 50.8 52.6 4122 10670 10649 51.3 40.9 0.5 744 75.7 40.9 35 53.4 603 1623 2427 2445 52.1 52.6 4123 3052 3033 50.3 50 1.8 626 76.4 42.8 35 53.6 604 1624 24418 24438 50 47.4 4124 24527 24507 51 42.9 1 110 71.3 42.7 35 50 605 1625 24417 24438 52.6 50.4 125 24517 24494 53.2 41.7 0.6 101 71.1 43.6 35 50.6 606 1626 8375 8396 51.8 45.5 4125 8298 9811 53.4 52.6 1.6 555 75.1 40 35 53.6 607 1627 24418 24439 52.9 45.5 4126 8929 8911 53.4 52.6 1.6 555 75.1 40 35 53.6 609 1628 8375 8396 51.8 45.5 4126 8929 8911 50.2 40.9 0.9 559 76.2 42.6 36 53.6 609 1628 818074 18094 51.1 42.9 4128 18662 18641 50.4 40.9 0.6 589 76.2 42.6 36 53.6 609 1629 18074 18094 51.1 42.9 4128 18662 18641 50.4 40.9 0.6 589 76.2 42.6 36 53.6 609 1629 18074 18094 51.1 42.9 4128 18662 18641 50.2 40.9 0.9 559 76.2 42.6 36 53.6 609 1629 18074 18094 51.1 42.9 4138 1888 8169 50.5 445 0.3 789 76.4 42.2 36 53.6 612 1632 3792 3811 54 55 4132 4446 4424 52.4 43.5 1.6 655 75.5 40.6 36 53.6 612 1632 3792 3811 54 55 4132 4446 4424 52.4 43.5 1.6 655 75.5 40.6 36 53.6 612 1632 3792 3811 54 55 4132 4446 4424 52.4 43.5 1.6 655 75.5 40.6 36 53.6 612 1632 3792 3811 54 55 4133 13545 13526 50.9 0.9 0.9 401 77.1 74.7 40.1 36 52.7 613 1633 25782 25805 52.1 41.7 4133 26182 26181 51.2 40.9 0.9 401 77.1 74.7 40.1 36 52.7 613 1633 25782 25805 52.1 41.7 4133 126182 26181 51.2 40.9 0.9 401 77.1 74.7 40.1 36 52.7 619 1638 2782 25805 52.1 41.7 4133 13565 13526 52.9 55 0.5 316 77.1 47.2 36 54.8 619 1638 25782 25805 52.1 41.7 4133 15618 26182 52.9 55 0.2 47.7 40.1 36 52.7 619 1639 13176 13197 52.7 45.5 4134 13545 13526 52.9 55 0.2 47.7 40.1 36 52.7 619 1639 13176 13199 52.7 45.5 4134 13545 13526 52.9 55 0.2 47.7 4.7 40.1 36 52.7 619 1638 25782 25805 52.1 47.7 4133 13565 13526 50.2 45.0 0.0 43.3 77.7 47.2 36 54.8 619 1638 25782 25805 52.1 47.7 4133 13565 13526 50.2 45.0 0.2 650 76.4 42.2 36 53.6 619 1638 25782 25806 52.9 55 0.4 137 13565 13526 50.2 45.0 0.0 43.3 76.3 42.2 36 53.6 619 1639	599	1619	7679	7698	50.6	50 4119	8049	8032	50.4	50	0.2	371	75.4	42.3	35	53
602 1622 9927 9945 50.8 52.6 4122 10670 10649 51.3 40.9 0.5 744 75.7 40.9 35 53.4 603 1623 2427 2445 52.1 52.6 4123 3052 3033 50.3 50 1.8 626 76.4 42.8 35 53.6 603 1625 24417 24436 55.0 47.4 4124 24527 24507 51 42.9 1 1 10 71.3 42.7 35 50.6 605 1625 24417 24438 55.0 4125 24517 24494 53.2 41.7 0.6 101 71.1 42.7 35 50.6 600 1626 8375 8396 51.8 45.5 4126 8929 8911 53.4 52.6 1.6 555 75.1 40.3 55 56 600 663 75.8 41.2 35 54 <	600	1620	13177	13197	50.3	42.9 4120	13320	13300	51.4	47.6	1.1	144	73.2	43.8	35	51.4
603 1623	601	1621	24179	24200	53.3	40.9 4121	24934	24913	53.4	45.5	0.2	.756	75.8	· 41	35	54.2
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605 1625 24417 24436 52.6 50 4125 24517 24494 53.2 41.7 0.6 101 71.1 43.6 35 50.6 606 1626 8375 8396 51.8 45.5 1472 8929 8911 53.4 52.6 1.6 555 75.1 40 35 53.2 607 1627 24418 24439 52.9 45.5 1472 25080 25082 53.5 52.6 0.6 663 75.8 41.2 35 54.6 608 1628 18074 18094 51.1 42.9 4129 18602 18641 50.2 40.9 0.9 569 76.2 42.8 36 53.6 610 1630 13231 13251 50.1 42.9 4130 13545 13527 50.3 52.8 0.2 315 74 47 47 46 53.6 61 1631 7400 7417 50.2	603	1623	2427	2445	52.1	52.6 4123	3052	3033	50.3	50	1.8	626	76.4	42.8	35	53.6
606 1626 8375 8396 51.8 45.5 4126 8929 8911 53.4 52.6 1.6 555 75.1 40 35 53.2 607 1627 24418 24439 52.9 45.5 4127 25000 25062 53.5 52.6 0.6 663 75.8 41.2 35 54 609 1629 18074 18094 51.1 42.9 4128 18682 18614 50.4 40.9 0.6 659 76.2 42.8 36 53.5 610 1630 13231 13251 50.1 42.9 4129 18632 18611 50.3 52.6 0.2 315 77 47 36 53.5 610 1630 13231 13251 50.1 42.9 4130 13527 50.3 52.6 0.2 315 77 47 43 55.4 611 1631 7401 7417 50.2	604	1624	24418	24436	50	47.4 4124	24527	24507	51	42.9	1	110	71.3	42.7	35	50
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608 1628 18074 18094 51.1 42.9 4128 18662 18641 50.4 40.9 0.6 589 76.2 42.6 36 53.6 609 1629 18074 18094 51.1 42.9 4129 18632 18811 50.2 40.9 0.6 589 76.2 42.6 36 53.6 610 1630 13231 13251 50.1 42.9 4130 13545 150.2 40.9 0.9 559 76.2 42.8 83.5 654 611 1631 7400 7417 50.2 50.0 4131 8188 8189 50.5 45 0.3 789 76.4 42.2 36 53.6 612 1632 3792 3811 5 554132 4444 4424 52.4 43.5 1.6 655 75.5 40.6 805 75.5 40.6 80.5 75.5 40.6 655 75.5 40.6 80.5 <t< td=""><td>606</td><td>1626</td><td></td><td>8396</td><td>51.8</td><td>45.5 4126</td><td>8929</td><td>8911</td><td>53.4</td><td>52.6</td><td>1.6</td><td>555</td><td>75.1</td><td>· 40</td><td>35</td><td>53.2</td></t<>	606	1626		8396	51.8	45.5 4126	8929	8911	53.4	52.6	1.6	555	75.1	· 40	35	53.2
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610 1830 13231 13251 50.1 42.9 4130 13545 13527 50.3 52.6 0.2 315 77 47 36 54 611 1831 7400 7417 50.2 50 4131 8188 8169 50.5 45 0.3 789 76.4 42.2 36 53.6 612 1832 3792 3811 54 55 4132 4446 4424 52.4 43.5 1.6 655 75.5 40.6 36 53.7 613 1833 25782 25805 52.1 41.7 4133 26182 26181 51.2 40.9 0.9 401 74.7 40.1 36 52.7 614 1834 13230 13251 52.4 45.5 4134 13545 13526 52.9 55 0.5 316 77.1 47.2 36 54.8 615 1835 985 1004 51.1 50 4135 1480 1482 51.6 47.4 0.5 486 76.4 43.5 36 52.7 616 1836 7400 7447 50.2 50 4136 8049 8032 50.4 50 0.2 860 76.1 42.2 36 53.6 617 1837 13176 13197 52.7 45.5 4137 13545 13526 52.9 55 0.2 370 77 46.2 36 54.8 618 1838 25782 25806 53.5 40 4138 26183 26182 52.9 55 0.2 370 77 46.2 36 54.8 619 1839 13176 13196 51.4 47.6 4139 13547 13526 52.9 55 0.2 370 77 46.2 36 54.8 619 1839 13176 13196 51.4 47.6 4139 13547 13526 52.9 55 0.2 370 77 46.2 36 54.8 619 1839 13176 13196 51.4 47.6 4139 13547 13526 50.2 45 1.2 372 76.9 46 36 54 6.2 1840 13298 132956 50.1 47.4 4140 13155 13138 50.4 50 0.3 218 75.4 45.4 36 52.9 621 1641 18080 18099 53 50 4141 18712 18693 54.8 55 1.9 633 76.3 42.7 36 54.8 622 1642 9140 9159 50.1 45 4142 9375 9364 50.4 40.9 0.3 218 75.4 45.4 36 52.9 623 1643 7725 7742 50 50 4136 8054 8035 50.4 50 0.4 30 37 75 41.8 36 52.9 625 1643 7725 7742 50 50 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12956 50.1 45.4 4140 13155 13138 50.4 50 0.3 218 75.4 45.4 36 52.9 625 1642 9140 9159 50.1 45 4142 9375 9364 50.4 40.9 0.3 288 75.3 42.8 36 52.9 625 1643 7725 7742 50 50 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12957 50.9 45 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12957 50.9 45 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12365 50.9 50 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12365 50.9 50 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12365 50.9 50 4144 10455 10435 50.5 42.9 0.8 534 75.3 40.6 36 52.9 625 1645 12338 12365 50.9 50 4144 10455 10435 50.5 42.9 0.8 53		1628		18094	51.1	42.9 4128		18641	50.4	40.9	0.6	589	76.2	42.6	36	53.6
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		9 1649	26039				6 4149	26183	2616	4 5	1 45	1.6	14	71.9	40.	7 36	50.8
	63		11540			4 5	0 4150	1172	1170	8 50.	4 45	0.1					
	63		12962	1298	0 50.	7 47.	4 4151	1354	1352	7 50.	3 52.6	0.4					
	63	2 1652	12961	1298	0 53.	2 5	0 4152	13545	1352								
	63	3 1653	9055	9079	9 52.	3 4	0 4153	9369								-	
	63	4 1654	12965	1298	5 51.			13545									
	63	5 1655	26039	26058	3 54	1 5		26693							45.4		
	63	6 1656	26039	26058	54		4156	26692							41.1		
	63	7 1657	26039	26058	3 54		4157	26688							41		
	63	B 1658	26039					26684						1	40.8		53.6
	63	9 1659	26039					26683							40.9		54.1
	64	1660	12965				4160	13545				1.4			40.9		53.8
	64	1 1661	26039			_	4161	26183		_		1.7	581	77.4	45.4	36	54.6
	64	1662	9055	9079			4162	9365			-	1.2	145		40.7	36	51.3
	643	1663	19795	19814				19922	19902			0.2		75.3	42.8		53.6
	644		12965	12988		_	4164	13545		_		0.4	128	72.2	. 43		50.7
	645		26040	26061			4165	26693	13526			1.2	581	77.4	45.4	36	55.1
	646		26040	26061	56.4		4166		26674			1.6	654	75.7	41.1	36	54.6
	647		26040	26061	56.4		4167	26693	26673			1.1	654	75.7	41.1	36	54.7
	648		26040	26061	56.4		4168	26690	26669			0.1	651	75.7	.: '41	36	55
	649	1	26040	26061	56.4			26685	26666	54.8	55	1.6	646	75.7	· 41	36	54.5
		1670	18011	18031	54.5		4169	26685	26665	55.3	52.4	1.1	646	75.7	41	36	54.7
-	651		7876	7895			4170	18443	18424	55.9	55	1.4	433	76.1	43.2	36	54.7
1	652		3230	3249	51.5		4171	8049	8032	50.4	50	1.2	174	73.2	· 42	36	51.5
ł	653		19795				4172	3646	3625	52	40.9	1.9	417	75.2	41.2	36	52.8
ł	654		12366	19814	50.4		4173	19920	19899	50.2	40.9	0.3	126	72.1	42.9	36	50.6
ł	655			12384	51.7		4174	12993	12975	51.4	47.4	0.3	628	76.5	. 43	36	54
ł		1676	19793	19814	54		4175	20544	20524	52.3	47.6	1.7	752	75.4	. 40	36	53.6
ŀ	657		12366	12384	51.7		4176	12911	12892	50.5	50	1.2	546	76.1	42.5	36	53.5
ŀ		1677	7728	7746	51.7		4177	8188	8168	50.4	42.9	1.3	461	75.6	41.9	36	53.1
ŀ		1678	26421	26441	51.5		4178	27084	27063	51.6	40.9	0.2	664	77.3	45	36	54.7
ŀ		1679	9929	9946	50		4179	10455	10434	51.1	40.9	1.1	527	75.3	40.6	36	52.8
ŀ		1680	26421	26441	51.5	42.9	4180	27083	27062	50.7	40.9	0.8	663	77.4	45.1	36	54.5
L	661	1681	12236	12256	51.2	42.9	4181	12999	12980	50.6	40	0.6	764	76.4	42.4	36	53.8
ļ	662	1682	26421	26441	51.5	42.9	4182	26694	26677	51.4	50	0	274	74.9	42.7	36	53
L	663	1683	9929	9946	50	50	4183	10183	10166	50.9	50	0.8	255	75.3	43.9	36	52.8
L		1684	12234	12252	50.6	47.4	4184	13000	12981	51.1	45	0.5	767	76.4	42.5	36	53.8
L		1685	8868	8889	50.4	40.9	4185	9254	9236	50.6	47.4	0.2	387	75	41.1	36	52.7
L		1686	9130	9150	51.3	42.9	4186	9597	9577	50.3	42.9	1	468	75.4	41.2	36	52.7
L	667	1687	9935	9955	50.4	42.9	4187	10605	10588	51.1	50	0.7	671	75.8	41.1	36	53.2
L		1688	26421	26441	51.5	42.9	4188	26587	26569	52	47.4	0.5	167	72.3			
L		1689	9130	9150	51.3	42.9		9597	9576	51	40.9	0.3	468	75.4	40.1	36	51.2
L	670	1690	26708	26727	50	45	4190	27466	27449	51	50	0.9	759		41.2	36	53.2
L	671	1691	9130	9150	51.3	42.9	4191	9375	9354	50.4	40.9	0.9	246	76	41.4	36	53.3
Γ	672	1692	10246	10266	50.4	47.6		10608	10589	51	50	0.5	363	74.7	42.7	36	52.5
Γ	673	1693	9924	9944	53.1	52.4		10449	10425	54.6	40	1.5	526	74.5	40.2	36	52.4
Γ	674	1694	12366	12384	51.7	52.6		12911	12891	51.2				75.4	40.9	36	53.8
Γ	675	1695	26708	26731	54.2	41.7		27466	27448	52.3	47.6	0.5	546	76.1	42.5	36	53.7
Γ	676	1696	8867	8888	52.7	45.5		9107	9086	51.6	52.6	1.9	759	76	41.4	36	54
۲	677	1697	9131	9151	50.4	42.9		9597			45.5	1.1	241	74.1	41.5	36	52.5
Γ	678	1698	9131	9151	50.4	42.9		9597	9577 9576	50.3	42.9	0.1	467	75.4	41.3	36	53
Γ		1699	10242	10265	51.2	41.7		10608	10589	51	40.9	0.6	467	75.4	41.3	36	53
				.0200	31.2	71.7	1138	10008	10589	51	50	0.3	367	74.5	40.1	36	52.5

B81 1701 27361 27380 52.4 55 4201 27467 27450 52.1 5.0 0.3 107 72.4 45.8 36 51.4 682 1702 27361 27380 52.4 55 4202 27466 27449 51 50 1.4 106 72.5 46.2 36 15.4 683 1703 3962 9944 50.5 52.6 4203 10449 10428 51.9 40.9 1.4 524 75.4 40.8 36 53 684 1704 9926 9944 50.5 52.6 4204 10449 10428 51.9 40.9 1.4 524 75.4 40.8 36 53 685 1705 13902 19802 53 52.6 4205 19922 19901 51.5 45.5 1.4 121 72.3 45.8 36 51.2 686 1706 27361 27380 52.4 55 4206 27482 27443 51.4 45 1 102 71.8 45.1 36 50.8 687 1707 1010 10159 52.4 50 4207 10605 16088 51.1 50 1.3 468 75 40.3 36 50.8 688 1708 16366 16384 52.9 55 4206 16777 16758 51.5 50 1.2 412 75.1 41 36 52.8 689 1709 16366 16385 52.9 55 4209 16781 16761 51.3 47.6 1.6 416 75.1 41.1 36 52.8 699 1711 16366 16385 52.9 55 4211 16777 16758 51.5 50 1.2 412 75.1 41.1 36 53.1 692 1711 16366 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41.1 36 53.1 692 1711 16366 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41.1 36 53.1 40.9 1711 16366 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41.1 36 53.1 40.9 1711 16761 1711																	
Fig. 2, 1702 27361 27380 52.4 55.4 202 27486 27449 51 50 1.4 106 72.5 46.2 30 51.1			27361	27380	52.4			27468	27451	51.1	50	1.3	108	72.3	45.4	36	51
683 1703 9926 9944 50.5 52.6 2603 10449 10428 51.9 40.9 1.4 524 75.4 40.8 36 53 685 1705 19802 19820 53 52.6 4204 10449 10431 50.9 47.4 6.5 524 75.4 40.8 36 53 685 1705 19802 19802 53 52.6 4206 19822 19801 51.5 45.5 41.1 12 71.8 43.8 36 51.2 686 1706 27361 27380 52.4 55 4206 27462 27443 51.4 45 1 102 71.8 45.1 36 50.8 687 1707 10140 10159 52.4 55 4206 27462 27443 51.4 45 1 102 71.8 45.1 36 50.8 688 1708 16386 16384 50.3 52.8 4208 16777 16758 51.5 50 1.2 412 75.1 41 36 52.8 688 1708 16386 16385 52.9 55 4209 16781 16761 51.3 467 47.6 1.8 416 75.1 41 36 52.8 689 1701 985 1008 56.1 50 4210 1484 1484 54.3 47.6 1.8 500 76.4 41.1 36 51.8 690 1711 16386 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41 36 52.8 691 1711 16386 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41 36 51.8 692 1712 27386 27344 52.4 41.5 41.3 462 54.3 47.6 1.8 40.9 76.4 43.5 36 54.8 692 1712 27386 27344 52.4 41.5 41.3 53.1 462 54.3 47.6 1.8 40.9 76.4 43.5 36 54.8 694 1714 2823 2844 50.4 45.5 4214 3052 30.3 50.3 50.0 2.2 207 74.1 41.7 5.2 566 1716 8867 8886 50.7 50 4216 3304 3485 50.4 45.0 50.4 44.7 4.2 36 52.5 689 1716 8867 8886 50.7 50 4216 3304 30.5 50.6 47.4 40.1 38 57.4 40.4 36 50.8 699 1718 8867 8886 50.7 50 4217 9244 9359 9368 51 50 0.6 63.5 75.8 421 36 53.8 699 1719 8867 8886 50.7 50 4217 63.4 63.6 50.6 47.6 40.9 60.9 74.1 40.9 50.8 699 74.1 40.9 50.8 699 74.1 40.9 60.9 74.1 40.9 60.9 74.1 40.9 74.1 40.9 60.9 74.1 40.9 74.1 4																36	51.4
684 1704 9926 9944 50.5 52.6 2204 10449 10431 50.9 47.4 0.5 524 75.4 40.6 36 53 686 1705 19802 19803 53 52.6 4205 19802 19801 51.5 45.5 1.4 121 72.3 43.8 36 51.2 586 1707 10140 10159 52.4 50.4207 10605 10588 51.1 50 1.3 466 75 40.3 36 52.6 688 1709 16386 16385 52.9 55 4209 16781 1677 16756 51.5 50 1.3 466 75 40.3 36 52.8 688 1709 16386 16385 52.9 55 4209 16781 16761 51.3 47.6 1.6 416 75.1 41.1 36 52.1 690 1710 985 1008 66.1 50 4210 1484 1464 54.3 47.6 1.6 416 75.1 41.1 36 52.1 690 1711 16386 16385 52.9 55 4211 16777 16756 51.5 50 1.4 412 75.1 41.1 36 52.1 690 1712 27386 27384 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.6 36 50.8 693 1713 986 1008 66.1 50 4213 1483 1462 54.3 45.5 18. 599 76.4 43.5 36 54.8 693 1713 986 1008 66.1 50 4213 1483 1462 54.3 45.5 18. 599 76.4 43.5 36 54.8 693 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50.0 2.20 77.4 41.7 41.3 65.1 696 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50.0 2.20 77.4 41.7 41.3 65.1 696 1716 3867 8886 50.7 50 4216 9399 9365 51.4 45.0 10.1 74.7 42.9 36 52.5 696 1716 3867 8886 50.7 50 4216 9390 9365 51.4 45.0 50.4 447 54.4 44.3 65.3 696 1719 8867 8887 52.3 47.6 4219 9369 9365 51.4 40.9 0.7 641 75.4 40.4 36 53.1 699 1719 8867 8887 52.3 47.6 4229 9341 9329 50.6 47.4 0.1 388 75.1 41.1 36 53.3 700 1721 9326 9346 50.4 42.2 42														72.5	46.2	36	51.1
686 1705 19802 19802 53 52.6 4205 19922 19901 51.5 45.5 1.4 121 72.3 43.6 36 51.2 686 1706 27381 27380 52.4 55 4206 27462 27443 51.4 45 1 102 71.8 45.1 36 50.8 687 1707 10140 10159 52.4 55 4208 10605 10586 51.1 50 1.3 466 75.1 40.3 36 52.9 688 1708 16386 16384 50.3 52.6 4208 16777 16756 51.5 50 1.2 412 75.1 41 36 52.8 689 1709 16386 16385 52.9 55 4210 1484 1484 54.3 47.6 1.6 416 75.1 41.1 36 52.8 691 1711 16366 16365 52.9 55 4211 16777 16756 51.5 50 1.4 412 75.1 41 36 53.1 692 1712 27386 27394 52.2 52.6 4212 27466 27446 2745 27386 27394 52.2 52.6 4212 27466 27446 2745 27386 27394 52.2 52.6 4212 27466 27446 2745 27386 27394 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.3 36 50.8 693 1713 985 1008 65.1 50 4213 1483 1462 54.3 45.5 1.8 499 76.4 43.5 36 54.8 695 1715 3224 3242 50.5 52.6 4215 3504 3465 50.4 45.5 1.8 499 76.4 43.5 36 54.8 696 1716 3824 3824 50.5 52.6 4215 3504 3465 50.6 47.5 0.5 444 75.4 41.4 36 52.6 696 1718 3839 3367 51.7 52.6 4218 9369 9968 51 40.3 0.7 641 75.4 40.4 36 52.5 696 1718 9349 3367 51.7 52.6 4218 9369 9968 51 40.3 0.7 641 75.4 40.4 36 52.8 699 1719 8867 8867 58.3 75.2 47.6 4219 9369 9360 51.5 50 0.5 638 76.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8199 8196 51 50.0 50.5 638 76.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8199 8196 51 50.0 40.5 67.6 76.8 74.1 93.6 52.4 70.5 70.1 70.1 70.1 70.2 70.2 70.7 70.1 70.5 70.4 70.5 70.4 70.5 70.4 70.5 70.4 70.5 70.4 70.5 70.5 70.5 70.5 70.5																36	
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687 1707 10140 10150 52.4 50 4207 10605 10586 51.1 50 1.3 466 75 40.3 56 52.9 688 1708 16386 16384 50.3 52.8 4208 16777 16756 51.5 50 1.2 412 75.1 41.3 65 52.8 689 1710 985 1008 56.1 50 4209 16781 16781 578 51.5 47.6 1.8 416 75.1 41.3 65 53.1 690 1710 985 1008 56.1 50 4210 1484 1464 54.3 47.6 1.8 500 76.4 43.6 36 54.9 691 1711 16386 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41.3 65 53.1 692 1712 27366 27384 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.6 36 53.1 692 1712 27366 27384 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.6 36 53.1 693 1713 985 1008 56.1 50 4213 1483 1462 54.3 45.5 1.8 499 76.4 43.5 36 54.8 694 1714 2823 2944 50.4 45.5 4214 3052 3033 50.3 50 0.2 220 71.4 14.7 47.7 42 36 52.5 695 1715 3224 3242 50.5 52.8 4215 3504 3485 50.4 45 0.1 281 74.7 42 36 52.5 696 1716 8667 888 50.7 50 4217 9254 9236 50.6 477.4 0.1 388 71.4 14.7 36 53.1 697 1717 8867 8885 50.7 50 4217 9254 9236 50.6 477.4 0.1 388 71.4 14.3 65 53.1 697 1717 8867 8887 52.3 47.8 4219 9369 9368 51.5 50 0.8 470 0.1 387 17.4 14.3 65 53.1 697 1719 8867 8887 52.3 47.8 4219 9369 9360 51.5 50 0.8 503 75.8 421 36 53.6 699 1719 8867 8887 52.3 47.8 4229 9341 9322 51.1 50 0.8 503 75.8 421 36 53.6 700 1720 8867 8887 52.3 47.8 4229 9341 9322 51.1 50 0.5 683 75.8 421 36 53.6 700 1720 8867 8887 52.3 47.8 4229 9349 349 349 349 349 349 349 349 349 3	685	1705	19802			52.6	4205	19922	19901	51.5	45.5	1.4	121	72.3	43.8	36	51.2
688 1708 16366 16384 50.3 52.6 4208 16777 16756 51.5 50 1.2 412 75.1 41 36 52.8 689 1709 16366 16385 52.9 55 4209 16781 16781 51.3 47.6 1.6 416 75.1 41.3 56 53.1 690 1710 995 1006 56.1 50 4210 1484 1464 54.3 47.6 1.6 1.8 500 76.4 43.6 3.6 54.9 691 1711 115366 16385 52.9 55 4211 16777 16758 51.5 50 1.4 412 75.1 41 36 53.1 692 1712 27366 27386 52.2 52.6 4212 27466 27446 52.3 52.6 0.1 101 71.5 44.0 36 50.8 693 1713 995 1006 56.1 50 4213 1483 1462 54.3 47.6 1.6 10 171.5 44.6 36 50.8 693 1713 985 1006 56.1 50 4213 1483 1462 54.3 47.6 1.6 10 171.5 44.6 36 50.8 694 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50 0.2 20 74.1 41.7 36 52.6 696 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 54.1 47.7 42 36 52.5 666 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 544 77.4 41.3 36 53.1 697 1717 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 544 74.4 0.1 388 75.1 41.2 36 52.5 699 1719 8867 8887 52.3 47.6 4220 9341 9322 51.1 50 0.8 50.3 75.8 42.1 36 53.6 699 1719 8867 8887 52.3 47.6 4220 9341 9322 51.1 50 0.2 475 75.7 4.1 36 53.4 701 1721 9926 9944 50.5 52.6 4221 10608 10689 15 0.0 50.5 683 175.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.4 40.9 0 245 74.7 42.3 65 53.5 704 1724 3055 3075 51.8 47.6 4223 3937 50.5 12 50.4 50.5 64.0 40.9 0 245 74.7 42.3 6 53.5 704 1724 3055 3075 51.8 47.6 4223 3937 50.8 1728 30.3 40.9 0 245 74.7 42.3 65 53.5 704 1724 3055 3075 51.8 47.6 4223 3937 50.5 1.5 50 0.8 503 75.8 42.1 36 53.4 70.1 1721 9926 9944 50.5 52.6 4221 10608 10689 15 50.0 5.0 583 175.8 41.1 36 53.4 70.1 1721 9926 9945 50.5 52.6 4221 10608 10689 51 50 0.5 683 75.8 41.8 36 53.4 70.1 1721 9926 9945 50.5 52.6 4221 10608 10689 51 50 0.5 683 75.8 41.1 36 53.4 70.1 1721 9926 9945 50.5 50.4 42.9 4223 3937 50.5 17.6 17.2 17.2 17.2 17.2 17.2 17.2 17.2 17.2	686	1706							27443		45	1			45.1	36	50.8
689 7709 16366 16385 52.9 55 4209 16781 16761 51.3 47.6 1.6 416 75.1 41.1 56 53.1 689 1711 16366 16385 52.9 55 4210 1484 1464 54.3 47.6 1.8 500 76.4 43.6 36 54.9 692 1712 27366 27384 52.2 55.2411 16777 16758 51.5 50.1 1.4 17.5 1.4 36 54.9 692 1712 27366 27384 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.6 36 50.8 693 1713 985 1008 56.1 50 4213 1483 1462 54.3 45.5 1.8 499 76.4 43.5 36 54.8 694 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50 0.2 230 74.1 41.7 36 52 695 1716 3824 3242 50.5 52.6 4215 3504 3485 50.4 45 0.5 447 47.5 41.4 36 52.5 696 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 447 47.5 41.4 36 52.5 697 1717 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 447 47.5 41.4 36 52.8 699 1719 8867 8886 50.7 50 4216 9369 9360 51.5 50 0.8 503 68 42.1 36 53.8 700 1720 8867 8887 52.3 47.6 4220 9341 9322 51.1 50 1.2 475 75.7 41.9 36 53.8 701 1721 9926 9944 50.5 52.6 4221 10608 10589 51.5 50 3.5 683 75.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 466 75.6 41.8 36 53.3 703 1723 9131 9151 50.4 42.9 4223 9375 9364 50.4 40.9 0.2 45 74.7 42.9 65.5 704 1724 3055 3076 51.8 47.6 4229 8309 308 50.5 50.0 60.6 465 75.6 41.8 36 53.1 706 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 50.6 0.3 766 75.6 41.8 36 53.1 706 1727 12370 12388 50.1 47.6 4228 3290 3189 50.5 50.0 60.6 465 75.6 41.8 36 53.1 701 7130 7233 936 51.8 47.6 4229 9369 350 50.8 50.0 60.8 40.5 75.6 41.9 36	687	1707	10140	10159	52.4	50	4207	10605	10588	51.1	50	1.3	466	75	40.3	36	52.9
690 1710 985 1008 56.1 50 4210 1484 1484 54.3 47.6 1.8 500 76.4 43.6 58 54.9 691 1711 16368 16385 52.9 55 4211 16777 16768 51.5 50 1.4 412 75.1 41 36 53.1 692 1712 27368 27384 52.2 52.6 4211 16777 16768 51.5 50 1.4 412 75.1 41 36 53.1 693 1713 985 1008 56.1 50 4213 1483 1482 54.3 45.5 1.8 499 76.4 43.5 36 50.8 691 1716 2823 2844 50.4 45.5 4214 3052 3033 50.3 50 40 0.2 230 74.1 41.7 36 52.8 695 1715 3224 3242 50.5 52.6 4214 3052 3033 50.3 50 40 0.2 230 74.1 41.7 36 52.5 696 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 444 75.4 41.4 36 53.1 697 1717 8867 8886 50.7 50 4217 9254 9236 50.6 47.4 0.1 388 75.1 41.2 36 52.5 699 1719 8867 8888 50.7 50 4217 9254 9236 50.6 47.4 0.1 388 75.1 41.2 36 53.2 699 1719 8867 8888 52.3 47.6 4220 9349 9369 51.5 50 0.8 603 75.8 42.1 36 53.2 700 1720 8867 8887 52.3 47.6 4220 9341 9322 51.1 50 0.8 603 75.8 42.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.0 3. 466 75.6 41.1 36 53.3 703 1723 9131 9151 50.4 42.9 4223 89375 9354 00.4 40.9 0.2 467.7 47.7 42.9 55.3 703 1729 139 13 1915 150.4 42.9 4223 9375 9354 50.4 40.9 0.7 40.7 47.7 42.9 55.3 703 1729 8867 8888 50.7 51.8 47.6 4224 3494 3473 50.4 40.9 1.4 440 76 43 36 53.3 703 1729 1319 151 50.4 42.9 4223 8190 8172 50.3 47.4 0.1 3.8 47.4 40.9 6.2 47.7 42.9 50 50 4222 8190 8172 50.3 47.4 0.1 3.4 40.9 76.4 41.3 36 53.3 703 1723 9131 9151 50.4 42.9 4223 9375 9354 50.4 40.9 0.2 47.4 40.9 6.2 47.7 42.9 36 53.5 703 1727 12370 12388 50.1 47.4 4227 13155 13138 50.4 50.0 1.2 67.7 47.7 42.9 50 50 4225 8189 8170 50.8 50.0 6.6 68.7 5.6 41.9 36 53.4 701 1731 14951 14975 52.2 40 4231 15150 1318 50.4 50.0 10.6 465 75.6 41.9 36 53.4 701 1732 836 50.7 51.8 47.6 4228 3093 3199 50.8 40.4 40.9 0.4 440 76 43.3 65.3 701 1731 14951 14975 52.2 40 4231 15164 1512 90.3 50.0 10.6 465 75.6 41.9 36 53.3 701 1733 14951 14975 52.2 40 4231 15164 1512 90.3 50.0 10.6 465 75.6 41.9 36 53.3 701 1733 14951 14975 52.2 40 4231 15164 1512 90.3 50.0 10.0 71.5 45 36 50.5 71.8 1438 16981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 53.6 711 1731	688	1708	16366			52.6	4208	16777	16758	51.5	50	1.2	412	75.1	41	36	52.8
691 1711 16366 16385 52.9 55 4211 16777 16756 51.5 50 1.4 412 75.1 41 36 53.1 692 1712 27366 27348 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.6 36 50.8 693 1713 965 1008 56.1 50 4213 1483 1482 54.3 45.5 1.8 499 76.4 43.5 36 50.8 694 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50 0.2 230 74.1 41.7 36 52 695 1715 3244 3242 50.5 52.6 4215 3504 3485 50.4 45 0.1 281 74.7 42 36 52.5 696 1716 8867 8886 50.7 50 4216 9310 921 51.2 45 0.5 44.7 40.1 388 75.1 41.2 36 52.8 696 1718 8867 8886 50.7 50 4216 9310 921 51.2 45 0.5 44.7 40.1 388 75.1 41.2 36 52.8 699 1719 8867 8887 52.3 47.6 4219 9399 9688 51 40.9 0.7 641 75.4 41.4 36 53.1 700 1720 8867 8887 52.3 47.6 4219 9399 9580 51.5 0.8 50 0.8 75.0 4216 36.9 17171 926 8867 8887 52.3 47.6 4219 9399 9580 51.5 0.8 60.7 50 0.8 60.7 74.0 1721 926 8867 8887 52.3 47.6 4219 9399 9580 51.5 0.8 60.7 50 0.8 60.7 75.0 4216 36.0 50.0 50.0 50.0 50.0 50.0 50.0 50.0 5	689	1709	16366	16385	52.9	55	4209	16781	16761	51.3	47.6	1.6	416	75.1	41.1	_36	53.1
692 1712 27366 27384 52.2 52.6 4212 27466 27448 52.3 52.6 0.1 101 71.5 44.6 36 50.8 631 713 985 1008 56.1 50 4213 1483 1462 54.3 45.5 1.8 499 76.4 43.5 36 54.8 694 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50.0.2 20.7 41. 14.7 36 52 695 1715 3224 3242 50.5 52.6 4216 30504 3485 50.4 45 0.1 281 74.7 42 36 52.5 696 1716 8867 8868 50.7 50 4217 9254 9236 50.6 47.4 0.1 381 74.7 42 36 52.5 696 1716 8867 886 50.7 50 4217 9254 9236 50.6 47.4 0.1 381 75.1 412 36 52.5 699 1719 8867 8887 52.3 47.6 4219 9399 9368 51 40.9 0.7 641 75.4 40.4 36 53.1 700 1720 8867 8887 52.3 47.6 4219 9399 9368 51 51 50 0.2 50 47.7 40.1 38 70.7 40.1 38 70.7 40.1 37.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 38 70.7 40.1 40.7 40.1 38 70.7 40.1 38 70.7 40.1 40.7 40.1 38 70.7 40.1 40.1 40.7 40.1 40.1 40.7 40.4 36 53.2 40.7 40.1 40.1 40.7 40.1 40.1 40.7 40.1 40.1 40.7 40.1 40.1 40.7 40.1 40.1 40.1 40.1 40.1 40.7 40.1 40.1 40.1 40.1 40.1 40.1 40.1 40.1	690	1710	985	1008	56.1	50	4210	1484	1464	54.3	47.6	1.8	500	76.4	43.6	36	54.9
693 1713 985 1008 56.1 50 4213 1483 1462 54.3 45.5 1.8 499 76.4 43.5 36 54.8 694 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50 0.2 220 74.1 41.7 42 36 52.5 696 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 444 75.4 41.4 36 53.1 697 1717 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 444 75.4 41.4 36 53.1 698 1718 9349 9367 51.7 52.6 4218 9389 9380 50.6 67.4 0.1 388 75.1 41.2 36 52.8 699 1719 8867 8887 52.3 47.6 4219 9369 9350 51.5 50 0.8 503 75.8 42.1 36 53.6 700 1720 9867 8887 52.3 47.6 4219 9369 9350 51.5 50 0.8 503 75.8 42.1 36 53.6 701 1721 9926 9944 60.5 52.6 4221 10608 10589 51 50 0.5 683 75.8 41.1 36 53.8 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 406 75.6 41.8 36 53.7 704 1724 3055 3075 51.8 47.6 4224 3494 3473 50.4 40.9 0 245 74.7 42.9 36 52.5 706 1725 7725 7742 50 50 4225 8189 8170 50.6 50 0.6 405 75.6 41.9 36 53.4 706 1727 12370 12388 50.1 47.6 4228 3209 3189 50.5 47.6 1.3 15.5 74.1 41.9 36 53.1 708 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 15.5 74.1 45.2 36 53.1 709 1729 8867 8887 52.3 47.6 4228 3209 3189 50.5 47.6 1.3 15.5 74.1 45.2 36 53.1 709 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 15.5 74.1 45.2 36 53.1 709 1729 8867 8887 52.3 47.6 4228 3209 3189 50.5 47.6 1.3 15.5 74.1 45.2 36 52.1 707 1727 12370 27367 51.4 42.8 3209 3189 50.5 47.6 1.3 15.5 74.1 45.2 36 52.1 710 1730 27367 57.8 42.1 57.6 42.2 32.2 33.1 33.8 50.6 40.0 30.3 33.7 47.6 43.8 53.3 710 1730	691	1711	16366	16385	52.9	55	4211	16777	16758	51.5	50	1.4	412	75.1	41	36	53.1
694 1714 2823 2844 50.4 45.5 4214 3052 3033 50.3 50 0.2 200 74.1 41.7 36 52.6 696 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 444 75.4 41.4 36 53.1 697 1717 8867 8886 50.7 50 4217 9254 9236 50.6 47.4 0.1 388 75.1 41.2 36 52.8 698 1718 9349 9367 51.7 52.6 4218 9399 9989 51 40.9 0.7 641 75.4 40.4 36 53.1 699 1719 8867 8887 52.3 47.6 4219 9369 9360 51.5 50 0.8 75.8 441 36 53.1 700 1720 8867 8887 52.3 47.6 4219 9369 9360 51.5 50 0.8 50.8 75.8 41.1 36 53.4 701 1721 9926 9944 50.5 52.6 4221 6008 10589 51 50 0.5 683 75.8 41.1 36 53.4 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.6 40.9 0.2 45 74.7 42.9 36 52.5 704 1723 9131 9151 50.4 42.9 4223 9375 9384 50.4 40.9 0 245 74.7 42.9 36 52.5 706 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 234 74.2 41.9 36 53.4 706 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 706 1728 3055 3075 51.8 47.6 4222 3058 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 709 1729 8867 8887 52.3 47.6 4222 3093 3198 50.5 57.6 63 75.8 41.1 36 53.3 710 1730 27367 27385 51.8 52.6 4230 27466 27446 52.3 52.6 0.9 100 71.6 45 36 53.4 711 1731 14951 14975 52.2 40.4231 15146 15129 50.3 50.1 64 75.6 41.8 36 53.3 719 1739 3796 3814 50.8 52.6 4230 4444 4424 50.6 42.9 0.1 52.1 75.9 42.2 36 53.3 719 1739 3796 3814 50.8 52.6 4230 4444 4444 4425 50.6 42.9 0.1 52.1 75.9 42.2 36 53.3 719 1731 3733 12234 12252 50.6 47.4 4233 12999 12990 1306 50.6 446 55.6 441 56.3 65.1 711 1731 4	692	1712	27366	27384	52.2	52.6	4212	27466	27448	52.3	52.6	0.1	101	71.5	44.6	36	50.8
696 1716	693	1713		1008	56.1	50	4213	1483	1462	54.3	45.5	1.8	499	76.4	43.5	36	54.8
696 1716 8867 8886 50.7 50 4216 9310 9291 51.2 45 0.5 444 75.4 41.4 36 53.1 697 1717 8867 8886 50.7 50 4217 9254 9236 50.6 47.4 0.1 388 75.1 41.2 36 52.8 699 1718 9349 9367 51.7 52.6 4218 9989 9988 51.4 40.9 0.7 641 75.4 40.4 36 53.2 699 1719 8867 8887 52.3 47.6 4219 9369 9350 51.5 50 0.8 503 75.8 42.1 36 53.6 700 1720 98667 8887 52.3 47.6 4219 9369 9350 51.5 50 0.8 503 75.8 42.1 36 53.6 701 1721 9926 9944 60.5 52.6 4221 10608 10589 51 50 0.5 683 75.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 406 75.6 41.8 36 53.3 703 1723 9131 9151 50.4 42.9 4223 9375 9354 50.4 40.9 0 245 74.7 42.9 36 52.5 704 1724 3055 3075 51.8 47.6 4224 3494 3473 50.4 40.9 1.4 40.7 643 36 53.4 706 1725 7725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.1 706 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 707 1727 12370 12388 50.1 47.4 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 708 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 8867 8887 52.3 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 710 1730 27367 27385 51.4 52.6 4230 27468 22.3 50.6 40.0 71.6 44.5 74.4 44.8 63.3 710 1730 27367 27385 51.4 52.6 4230 27468 22.3 52.6 0.9 0.7 50 1.6 44.5 75.4 41.6 36 53.3 710 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 196 73.2 40.8 36.3 711 1731 1328 3867 3887 52.3 47.6 4230 27468 27546 27527 51.3 50	694	1714	2823	2844	50.4	45,5	4214	3052	3033	50.3	50	0.2	230	74.1	41.7	36	-52
697 1717	695	1715	3224	3242	50.5	52.6	4215	3504	3485	50.4	45	0.1	. 281	74.7	, 42	36	52.5
698 1718 9349 9367 51.7 52.6 4218 9989 9968 51 40.9 0.7 641 75.4 40.4 36 53.2 699 1719 8867 8887 52.3 47.6 4219 9369 9350 51.5 50 0.8 503 57.8 42.1 36 53.6 700 1720 8867 8887 52.3 47.6 4220 9341 9322 51.1 50 1.2 475 75.7 41.9 36 53.4 701 1721 9926 9944 50.5 52.6 4221 10608 10589 51 50 0.5 683 75.8 42.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 466 75.6 41.8 36 53.3 703 1723 9131 9151 50.4 42.9 4223 9375 9354 50.4 40.9 0 245 74.7 42.9 36 52.5 704 1724 3055 3075 51.8 47.6 4224 3494 3473 50.4 40.9 0 245 74.7 42.9 36 52.5 705 1725 7742 50 50 4228 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.4 706 1726 2623 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 707 1727 12370 12388 50.1 47.4 4227 3155 3138 50.4 50.0 3.7 786 76.8 43.4 36 53.9 708 1728 8667 8887 52.3 47.6 4229 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 8867 8867 52.3 47.6 4229 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 53.3 710 1730 27367 27385 51.4 52.6 4220 27466 27448 52.3 52.6 0.9 100 71.6 41.8 36 53.3 714 1734 3055 3076 52.4 45.5 4232 3049 319 50.8 45.5 1.6 477 57.4 41.9 36 53.3 715 1735 8867 8887 52.3 47.8 4232 3299 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 719 1730 27367 27385 51.4 52.6 4230 27466 27448 52.3 52.6 0.9 100 71.6 45.8 53.6 719 1733 12234 12252 50.6 47.4 4233 12999 12980 50.6 40 0 766 76.4 42.4 36 53.8 711 1731 34951 34951 3495 347.8 4235 347.8 4235 347.8 4235 347.8 4235 347.8 4235 347.8 4235 347.8	696	1716	8867	8886	50.7	50	4216	9310	9291	51.2		0.5	444	75.4	41.4	36	53.1
699 1719 8867 8887 52.3 47.6 4219 9369 9350 51.5 50 0.8 50.3 75.8 42.1 36 53.6 700 1720 98667 8887 52.3 47.6 4221 9369 9341 9322 51.1 50 0.5 683 75.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 466 75.6 41.8 36 53.7 703 1723 9131 9151 50.4 42.9 4223 8395 9354 50.4 40.9 0 245 74.7 42.9 36 52.5 703 1724 3055 3075 51.8 47.6 4224 3494 3473 50.4 40.9 10 245 74.7 42.9 36 52.5 705 1725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.1 706 1725 7725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.1 706 1725 7725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.1 706 1725 2238 50.1 47.4 4227 3055 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 707 1727 12370 12388 50.1 47.4 4227 3155 3138 50.4 50 0.3 766 76.8 43.4 38 53.3 706 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 2867 8887 52.3 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 710 1730 27367 27385 51.4 52.6 4230 27468 52.3 52.6 0.9 100 71.6 44.8 53.3 710 1730 27367 27385 51.4 52.6 4230 27468 52.3 52.6 0.9 100 71.6 44.8 53.6 53.1 711 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 196 73.2 40.8 36 51.4 711 7133 12234 12252 50.6 47.4 4233 12399 12990 50.6 40 0.7 67.6 76.6 42.4 36 53.8 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 366 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 717 1737 2671 2692 52.1 40.9	697	1717	8867	8886	50.7	50	4217	9254	9236	50.6	47.4	0.1	388	75.1	41.2	36	52.8
700 1720 8867 8887 52.3 47.6 4220 9341 9322 51.1 50 1.2 475 75.7 41.9 36 53.4 701 1721 9926 9944 50.5 52.6 4221 10508 10589 51 50 0.5 683 75.8 41.1 36 53.3 702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 466 75.6 41.1 36 53.3 703 1723 9131 9151 50.4 42.9 4223 9375 9354 50.4 40.9 0 245 74.7 42.9 36 52.5 704 1724 3055 3075 51.8 47.6 4224 3949 3473 50.4 40.9 0 245 74.7 42.9 36 52.5 704 1724 3055 3075 51.8 47.6 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.4 705 1725 7725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.4 706 1726 2823 2844 50.4 45.5 4226 3056 3036 50.8 52.6 0.3 234 74.2 41.9 36 52.2 707 1727 12370 12388 50.1 47.6 4225 3193 81.0 50.6 50 0.6 465 75.6 41.9 36 53.4 706 1728 3055 3075 51.8 47.6 4229 9340 9319 50.8 47.6 42.9 13.1 15.5 13.1 15	698	1718	9349	9367	51.7	52.6	4218	9989	9968	51	40.9	0.7	641	75.4	40.4	36	53.2
Total Tota	699	1719	8867	8887	52.3	47.6	4219	9369	9350	51.5	50	0.8	. 203	75.8	42.1	36	53.6
702 1722 7725 7742 50 50 4222 8190 8172 50.3 47.4 0.3 466 75.6 41.8 36 53 703 1723 9131 9151 50.4 42.9 4223 9375 9354 50.4 40.9 0 245 74.7 42.9 36 52.5 704 1724 3055 3075 51.8 47.6 4224 3494 3473 50.4 40.9 1.0 440 76 43.9 56 52.5 705 1725 7725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.1 706 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 707 1727 12370 12388 50.1 47.4 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 8867 8887 52.3 47.6 4229 9340 9319 50.8 45.5 1.6 474 75.6 41.8 36 53.3 710 1730 27367 27385 51.4 52.6 4230 27466 2748 52.3 52.6 0.9 1.9 196 73.2 40.8 36 53.4 711 1731 14951 14975 52.2 40 4231 15154 15129 50.3 50 1.9 196 73.2 40.8 36 53.4 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4232 9311 9290 50.7 50 1.6 445 75.4 41.6 36 53.1 716 1739 3056 3076 52.4 45.5 4236 3495 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4236 329 311 9290 50.7 50 1.6 445 75.4 41.6 36 53.1 716 1739 1739 1739 1739 1739 1739 1739 1739	700	1720	8867	8887	52.3	47.6	4220	9341	9322	51.1	50	1.2	475	75.7	41.9	36	53.4
703 1723 9131 9151 50.4 42.9 4223 9375 9384 50.4 40.9 0 245 74.7 42.9 36 52.5 704 1724 3055 3075 51.8 47.6 422 3194 3473 50.4 40.9 1.4 40 76 43 36 53.4 706 1725 7725 50 50 50 62.5 60 0.6 465 75.6 41.9 36 53.4 706 1726 2823 2844 50.4 45.5 4226 3056 3036 50.8 52.6 0.3 234 74.2 41.9 36 52.2 706 1728 3055 3075 51.8 47.6 4229 3040 9319 50.5 47.6 41.8 36 59.3 71.0 71.6 46 52 36.6 50.0 30.1 71.1 71.6 45 36.8 53.1	701	1721	9926	9944	50.5	52.6	4221	10608	10589	51	50	0.5	683	75.8	41.1	36	53.3
704 1724 3055 3075 51.8 47.6 4224 3494 3473 50.4 40.9 1.4 440 76 43 36 53.4 706 1725 7742 50 50 4225 8189 8170 50.6 50 0.6 465 75.6 41.9 36 53.1 706 1728 2823 2844 50.4 45.5 4227 13155 13138 50.4 50 0.3 786 76.8 43.9 36 52.2 700 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 700 1729 8867 8887 52.3 47.6 4229 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 70.1 1731 14951 14951 52.2 40.4231 <	702	1722	7725	7742	50	50	4222	8190	8172	50.3	47.4	0.3	. 466	75.6	41.8	36	53
705 1725 77725 77742 50 50 4225 8189 8170 50.6 50 0.6 405 75.6 41.9 36 53.1 706 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 234 74.2 41.9 36 52.2 707 1727 12370 12389 50.1 47.4 4227 13155 13138 50.4 50 0.3 766 76.8 43.4 38 53.3 708 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 8867 8887 52.3 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 710 1730 27367 27385 51.4 52.6 4220 27468 52.3 52.6 0.9 1729 8867 8887 52.3 47.6 4229 9340 9319 50.8 45.5 1.6 474 75.6 41.8 36 53.3 710 1730 27367 27385 51.4 52.6 4230 27466 27486 52.3 52.6 0.9 1729 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 196 73.2 40.8 36 51.4 712 1732 8867 8887 52.3 47.6 4229 2931 9292 50.7 50 1.6 445 75.4 41.6 36 53.1 712 1732 8867 8887 52.3 47.6 4228 29311 9292 50.7 50 1.6 445 75.4 41.6 36 53.1 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4236 59109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 3055 3076 52.4 45.5 4236 3209 3189 50.5 47.6 2.1 55 74.1 42.2 36 52.1 717 1737 2671 2692 52.1 40.9 4237 3053 3034 50.3 50 1.8 383 74.7 40.5 36 52.5 718 1738 16981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 53.6 719 1739 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.1 521 75.9 42.2 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3798 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3798 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3798 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3798 3814 50.8 52.6 4240 4445 4225 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3798 3814 50.8 52.6 4240 4445 4225 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1743 27383 27403	703	1723	9131	9151	50.4	42.9	4223	9375	9354	50.4	40.9	0	245	74.7	42.9	36	52.5
766 1726 2823 2844 50.4 45.5 4226 3056 3038 50.8 52.6 0.3 224 74.2 41.9 36 52.2 707 1727 12388 50.1 47.4 4227 13156 13138 50.4 50 0.3 786 76.8 43.4 38 53.9 708 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 8667 8887 52.3 47.6 4228 9340 9319 50.8 45.5 1.6 474 75.6 41.8 36 53.3 710 1730 27367 27385 51.4 52.6 4209 27466 27448 52.3 50.1 19 17.16 45 50.6 40.7 17.11 1731 14951 14951 1429 50.7 50 <td>704</td> <td>1724</td> <td>3055</td> <td>3075</td> <td>51.8</td> <td>47.6</td> <td>4224</td> <td>3494</td> <td>3473</td> <td>50.4</td> <td>40.9</td> <td>1.4</td> <td>440</td> <td>76</td> <td>43</td> <td>36</td> <td>53.4</td>	704	1724	3055	3075	51.8	47.6	4224	3494	3473	50.4	40.9	1.4	440	76	43	36	53.4
707 1727 12370 12388 50.1 47.4 4227 13155 13138 50.4 50 0.3 768 76.8 43.4 36 53.9 708 1728 3055 3075 51.8 47.6 4229 3209 3189 50.5 47.6 1.3 155 74.1 45.2 38 52.1 709 1729 8867 8867 887 52.3 47.6 4229 9340 9319 50.8 45.5 1.6 474 75.6 44.8 36 53.3 710 1730 27367 27385 51.4 52.6 4230 27466 27448 52.3 52.6 0.9 100 71.6 -45 36 50.6 711 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 198 73.2 40.8 85.1 712 1732 8867 8887 52.3 47.6 40.6 6.6 40 1.	705	1725	7725	7742	50	50	4225	8189	8170	50.6	50	0.6	465	75.6	41.9	36	53.1
708 1728 3055 3075 51.8 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 709 1729 3867 8867 8887 52.3 47.6 4228 3209 3189 50.5 47.6 1.3 155 74.1 45.2 36 52.1 710 1730 27367 27385 51.4 52.6 4230 27466 27527 51.3 50 1.6 474 75.5 41.8 36 53.3 711 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 196 73.2 40.8 36 51.4 712 1732 8867 8887 52.3 47.6 4228 29311 9292 50.7 50 1.6 445 75.4 41.6 36 53.4 713 1733 12234 12252 50.6 47.4 4233 12999 12990 50.6 40 0 76 76.4 42.4 36 53.8 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 3667 3655 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 716 1736 3055 3076 52.4 45.5 4236 3209 3189 50.5 47.6 22 51.5 74.1 42.2 36 52.1 717 1737 2671 2692 52.1 40.9 4237 3053 3034 50.3 50 1.8 383 74.7 40.5 36 52.5 718 1738 18981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 721 1741 27382 27401 50.8 45 4241 27546 27527 51.3 50 0.6 165 73.7 43.6 36 51.7 722 1743 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.7 723 1743 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.7 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.7 726 1748 1779 17811 52.9 43.5 4246 18220 18202 54.8 52.6 0.9 230 74.9 40.5 36 53.4 727 1747 18004 18023 51.1 50 4247 18233 18215 51.3 50 1.1 10 72	706	1726	2823	2844	50.4	45.5	4226	3056	3038	50.8	52.6	0.3	234	74.2	41.9	36	52.2
709 1729 8867 8887 52.3 47.6 4229 9340 9319 50.8 45.5 1.6 474 75.6 41.8 36 53.3 710 1730 27367 27385 51.4 52.6 4230 27446 27446 27446 52.3 52.6 0.9 100 71.6 45 36 50.6 711 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 136 73.2 40.8 36 51.4 712 1732 8867 8887 52.3 47.6 4232 9311 9292 50.7 50 1.6 445 75.4 41.6 36 53.1 713 1733 12234 12252 50.6 47.4 4233 12999 12980 50.6 40 0 766 76.4 42.4 36 53.4 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4255 9109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 3055 3076 52.4 45.5 4236 3209 3189 50.5 47.6 2 155 74.1 45.2 36 52.1 717 1737 2671 2689 52.1 40.9 4237 3053 3034 50.3 50 1.8 383 74.7 40.5 36 52.5 719 1739 3796 3814 50.8 52.6 4239 4444 4424 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4239 4444 4424 50.6 42.9 0.2 650 75.5 40.5 36 53.1 721 1741 27382 27401 50.8 45.4241 27546 27527 51.3 50 0.6 160 73.4 43.1 36 51.5 723 1743 27383 27403 50.3 42.9 4244 27546 27527 51.3 50 1.1 164 73.5 43.3 36 51.5 726 1746 17791 17813 52.9 43.5 4245 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 726 1748 16004 18023 51.1 50.4247 18233 18215 51.3 50 0.2 20 75 40.5 36 53.4 728 1748 18004 18023 51.1 50.4247 18233 18215 51.3 50 0.1 110 72.4 45.5 36 50.8 729 1749 27437 27456 50.2 40.4249 27546 27527 51.3 50 1.1 110 72.4 45.5 36 50.8 729 1748 8004 18023 51.1 50.4247 18233 18215 51.3 50.6 1.1 128 74.7 40.5 36 53.4	707	1727	12370	12388	50.1	47.4	4227	13155	13138	50.4	50	0.3	786	76.8	43.4	36	53.9
710 1730	708	1728	3055	3075	51.8	47.6	4228	3209	3189	50.5	47.6	1.3	155	74.1	45.2	36	52.1
711 1731 14951 14975 52.2 40 4231 15146 15129 50.3 50 1.9 196 73.2 40.8 36 51.4 7121732 8667 8887 52.3 47.6 4252 9311 9292 50.7 50 1.6 445 75.4 41.6 36 53.1 7131733 12234 12252 50.6 47.4 4233 12999 12990 50.6 40 0 766 76.4 42.4 38 53.8 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8667 8887 52.3 47.6 4255 9109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8667 86.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8667 8887 52.3 47.6 425 9109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 3055 3076 52.4 45.5 4234 3495 3209 3189 50.5 47.6 22 155 74.1 452.3 6 52.1 717 1737 2671 2692 52.1 40.9 4237 3053 3034 50.3 50 1.8 383 74.7 40.5 36 52.5 718 1738 16981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 52.1 719 1739 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 721 1741 27382 27401 50.8 454241 27546 27527 51.3 50 1.6 16 160 73.4 43.1 36 51.5 724 1742 27382 27401 50.8 454241 27546 27527 51.3 50 1.6 16 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4243 27546 27527 51.3 50 1.1 164 73.5 43.3 36 51.7 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.4 725 1743 17769 17811 32.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 725 1744 17769 17813 52.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 725 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.4 725 1744 1744 27383 27403 50.3 42.9 4248 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.4 725 1744 1744 27383 27403 50.	709	1729	8867	8887	52.3	47.6	4229	9340	9319	50.8	45.5	1.6	474	75.6	41.8	36	53.3
712 1732 8867 8887 52.3 47.6 4232 9311 9292 50.7 50 1.6 445 75.4 41.6 36 53.1 713 1733 12234 12252 50.6 47.4 4233 12999 12990 50.6 40 0 766 76.4 42.4 36 53.8 714 1734 30.55 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4255 9109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 30.55 30.55 30.6 52.4 45.5 4236 3209 3189 50.5 47.6 2 155 74.1 45.2 36 52.1 716 1736 30.55 30.5 30.6 52.4 40.9 4237 30.53 30.34 50.3 50.1 8 333 74.7 40.5 36 52.1 717 1737 2671 2692 52.1 40.9 4237 30.53 30.34 50.3 50 1.8 333 74.7 40.5 36 52.5 718 1738 16981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 53.6 719 1739 3796 3814 50.8 52.6 4239 4444 4424 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4239 4444 4424 50.6 42.9 0.2 649 75.5 40.5 36 53.1 721 1741 27382 27401 50.8 45 4241 27546 27527 51.3 50 0.6 165 73.7 43.6 36 51.5 723 1743 27383 27403 50.3 42.9 4244 27546 27527 51.3 50 1.1 164 73.5 43.3 36 51.5 728 1745 27383 27403 50.3 42.9 4244 27546 27527 51.3 50 1.1 164 73.5 43.3 36 51.5 728 1745 17789 17811 52.9 43.5 4245 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 728 1748 18004 18023 51.1 50.4247 18233 18215 51.3 52.6 0.2 20 75 43.9 94.5 36 52.8 728 1748 18004 18023 51.1 50.4248 18231 18210 52.2 45.5 1.1 228 74.9 40.5 36 53.4 728 1748 18004 18023 51.1 50.4248 18231 18210 52.2 45.5 1.1 10 10 72.4 45.5 36 50.8 729 1748 18004 18023 51.1 50.4248 18231 18210 52.2 45.5 1.1 10 10 72.4 45.5 36 50.8 729 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 36 50.8 729 1748 18004 18023 51.1 50.4248 18231 18215 51.3 52.6 0.2 20 75 43.9 96 52.8 729 1748 18004 18023 51.1 50.4248 18231 18215 51.3 52.6 0.2 20 75 43.9 96 52.8 729 1748 18004 18023 51.1 50.4248 18231 18215 51.3 52.6 4.5 51.1 1228 74.7 40.5 36 53.4 729 1748 18004 18023 51.1 50.4248 18231 18215 51.3 50.1 1.1 110 72.4 45.5 36 50.8 729 1748 18004 18023 51.1 50.4248 18231 18215 51.3 50.1 1.1 110 72.4 45.5 36 50.8 729 1748 18004 18023 51.1 50.4248 18231 18215 51.3 50.1 1.1 110 72.4 45.5 36 50.8 729				27385				27466	27448	52.3	52.6	0.9	100	71.6	45	36	50.6
713 1733 12234 12252 50.6 47.4 4233 12999 12980 50.6 40 0 766 76.4 42.4 36 53.8 714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4235 9109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 3055 3076 52.4 45.5 4236 3209 3189 50.5 47.6 2 15.5 74.1 45.2 36 52.1 717 1737 2671 2692 52.1 40.9 4237 3053 3034 50.3 50 1.8 383 74.7 40.5 36 52.5 718 1738 16981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 53.6 719 1739 3796 3814 50.8 52.6 4239 4444 4424 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 649 75.5 40.5 36 53.1 721 1741 27382 27401 50.8 45 4241 27546 27527 51.3 50 0.6 165 73.7 43.6 36 51.9 722 1742 27382 27401 50.8 45 4242 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 723 1743 27383 27403 50.3 42.9 4243 27546 27527 51.3 50 0.1 169 73.2 42.8 36 51.4 725 1744 17784 17784 17789 1781 52.9 43.5 4245 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 726 1746 17789 17813 52.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 726 1746 17789 17813 52.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 728 1748 18004 18023 51.1 50 4247 18233 18215 51.3 52.6 0.2 230 75 43.9 36 52.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 36 53.4 726 1748 17789 17813 52.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 726 1748 17789 17813 52.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 728 1749 1748 18004 18023 51.1 50 4247 18233 18215 51.3 52.6 0.2 230 75 43.9 36 52.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 36 50.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 58 50.8 50.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 58 50.8 50.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 58 50.8 50.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 58 50.8 50.8 728 1749 27437 27456 50.2 40 4249 27546 27527 51.3 50 1.1 110 72.4 45.5 5											50	1.9	196	73.2	: 40.8	36	51.4
714 1734 3055 3076 52.4 45.5 4234 3495 3473 51.8 43.5 0.6 441 76 43.1 36 53.9 715 1735 8867 8887 52.3 47.6 4235 9109 9087 50.5 43.5 1.8 243 74 41.2 36 52.1 716 1736 3055 3076 52.4 45.5 4238 3209 3189 50.5 47.6 22 155 74.1 42.2 36 52.1 717 1737 2671 2692 52.1 40.9 4237 3053 3034 50.3 50 1.8 383 74.7 40.5 36 52.5 718 1738 16981 17000 51.3 50 4238 17501 17481 51.2 42.9 0.1 521 75.9 42.2 36 53.6 719 1739 3796 3814 50.8 52.6 4240 4445 4424 50.6 42.9 0.2 649 75.5 40.5 36 53.1 720 1740 3796 3814 50.8 52.6 4240 4445 4425 50.6 42.9 0.2 650 75.5 40.5 36 53.1 721 1741 27382 27401 50.8 454241 27546 27527 51.3 50 0.6 165 73.7 43.6 36 51.9 722 1742 27382 27401 50.8 454242 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4244 27546 27527 51.3 50 1.1 164 73.5 43.3 36 51.7 724 1744 27383 27403 50.3 42.9 4244 27546 27527 51.3 50 1.1 164 73.5 43.3 36 51.7 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.6 160 73.4 43.1 36 51.5 724 1744 27383 27403 50.3 42.9 4244 27541 27522 50.1 45 0.1 159 73.2 42.8 36 51.4 725 1743 17789 17811 32.9 43.5 4246 18220 18202 54.8 52.6 1.9 432 74.9 40.5 36 53.4 721 1747 18004 18023 51.1 50 4247 18233 18215 51.3 50 1.1 164 73.5 43.9 36 51.4 726 1748 17789 17811 32.9 43.5 4246 18220 18202 54.8 52.6 1.9 430 74.9 40.5 36 53.4 721 1747 18004 18023 51.1 50 4247 18233 18215 51.3 50 1.1 110 172.4 45.5 36 53.8 721 1749 27437 2745 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 36 53.8 722 1749 27457 2745 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 56 53.8 52.8 722 1749 27437 2745 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 56 53.8 52.8 722 1749 27457 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 56 53.8 52.8 722 1749 27457 2745 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 58 50.8 50.8 722 1749 27437 27455 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 58 50.8 50.8 722 1749 27437 27455 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 58 50.8 50.8 722 1749 27437 27455 50.2 40 4249 27546 27527 51.3 50 1.1 110 172.4 45.5 58 50.8 50.8 50.			8867		52.3			9311	9292	50.7	50	1.6	445	75.4	41.6	36	53.1
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	728	1748			51.1	50	4248			52.2		1.1		74.7	43.4	36	52.8
730 1750 27437 27456 50.2 40 4250 27541 27522 50.1 45 0.1 105 71.8 44.8 36 50.4															45.5	36	50.8
	730	1750	27437	27456	50.2	40	4250	27541	27522	50.1	45	0.1	105	71.8	44.8	36	50.4

732 1752 12233 12251 51.1 52.6 4252 12994 12976 50.3 47.4 0.8 762 76.5 42.5 36 53.7				-													
Total Tota						_		18223	18206	51.8	50	0.6	220	74.5	43.2	36	52.6
733 1755 7866 7869 52.5 47.6 4253 38192 3172 50.9 42.9 1.6 324 75.2 42.3 36 5.7 734 1755 3224 3242 50.5 52.6 4255 3500 3484 51.5 50 0.6 277 74.6 42.1 36 52.7 736 1756 3224 3242 50.5 52.6 4255 3500 3481 51.2 50 0.6 277 74.6 42 36 52.7 737 1757 1 22 54.8 50.4 4255 3497 3478 51.3 50 0.8 274 74.6 42 36 52.7 738 1759 9140 9159 50.1 45 4250 9597 9576 51 40.9 0.9 458 75.2 41.3 36 52.5 740 1760 9140 9159 50.1 45 4250 9597 9576 51 40.9 0.9 458 75.2 41.3 36 52.5 741 1761 7728 7746 51.5 52.6 4261 8188 8170 50.6 50.1 42.9 1.5 42.1 75.2 41.3 36 52.5 741 1761 7728 7746 51.5 52.6 4261 8188 8170 50.6 50.1 42.9 1.5 42.1 75.2 41.3 36 52.5 743 1763 12235 12243 50.1 52.6 4263 12999 12979 50.1 45 42.0 50.5 40.7 75.7 41.4 36 52.5 744 1766 3225 3244 52.4 52.4 52.4 52.4 53.4 50.0 3484 51.5 50.8 45 1.3 46.5 75.7 42.3 36 53.7 749 1768 3225 3244 52.4 55 4266 3500 3481 51.2 50.1 37.7 74.7 42.3 36 53.7 749 1769 12233 12235 15.1 52.6 4267 12999 12979 50.6 40 0.6 76.7 74.4 42.3 36 53.7 749 1769 12233 1225 15.1 52.6 4267 12999 12970 50.6 40 0.6 76.7 74.7 42.3 36 53.7 749 1769 12233 1225 15.1 52.6 4268 3907 3478 51.3 50 1.1 273 74.7 42.1 36 52.8 750 1777 12330 12251 51.1 52.6 4267 12999 12960 50.6 40 0.6 76.7 74.7 42.3 36 55.5 750 1777 12330 12251 51.1 52.6 4267 12999 12960 50.6 40 0.6 76.7 74.0 42.5 36 53.8 750 1777 12330 12251 51.1 52.6 4267 12999 12960 50.6 40 0.6 76.7 74.7 42.3 36 52.5 750 1777 13836 12414 51.5 54 4											47.4	8.0	762				53.7
734 1754 3224 3242 50.5 52.6 4255 3500 3481 51.5 50 0.9 280 74.8 42.1 56 52.1 756 1756 3224 3242 50.5 52.6 4255 3497 3476 51.3 50 0.8 277 74.6 41.9 36 52.1 776 11 22 54.8 50 4257 204 165 56.6 55 1.8 204 75.1 45.1 36 52.1 758 1758		1					4253	8192	8172	50.9	42.9	1.6	324	75.2	42.3	36	53
786 1756 3224 3242 50.5 52.6 4256 3500 3481 51.2 50 0.6 277 74.6 41.9 36 52.2 737 1757 11 22 54.8 50 4257 204 185 56.6 55 1.8 204 751 45.1 36 52.1 738 1758 9140 9159 50.1 45 4258 9597 9576 51 40.9 0.9 456 75.3 41.3 36 52.5 739 1759 28179 22800 50.8 40.9 4259 50.1 45.1 4258		1		3242	50.5	52.6	4254	3503	3484	51.5	50	0.9	280	74.8	42.1	36	
788 1756 3224 342 50.5 52.6 4256 3497 3478 51.3 50 0.8 274 74.6 42 36 52.2 733 1757 1 22 54.8 50.4257 20.4 185 56.6 55 18. 940 75.1 45.1 36 52.2 738 1759 28179 28200 50.8 40.9 4258 9867 9876 51 40.9 0.9 458 75.3 41.3 36 52.2 740 1760 9140 9159 50.1 45 4258 9867 28683 50.2 52.6 0.6 493 79.7 51.7 36 52.7 40 1760 9140 9159 50.1 45 42690 9960 540 51.6 42.9 1.5 421 75.7 42 36 52.2 741 1761 7728 7746 51.7 52.6 4261 8189 8170 50.6 50 1.1 482 75.7 42 36 52.2 742 1763 12235 12253 50.1 52.6 4263 12998 12979 50.1 45 0.7 64 76.3 41.4 36 52.2 744 1764 3225 3244 52.4 55 4264 3503 3444 51.5 50 1 279 74.8 42.3 36 52.2 745 1766 3225 3244 52.4 55 4264 3503 3444 51.5 50 1 279 74.8 42.3 36 52.2 747 1767 12233 12251 51.1 52.6 4267 12989 12979 50.8 45 13.3 46 75.5 40.6 32.2 344 52.4 55 4268 3500 3484 51.5 50 1 279 74.8 42.3 36 52.2 747 1767 12233 12251 51.1 52.6 4267 12989 12979 50.8 45 13.3 46 75.5 40.6 36 53.2 474 1767 12233 12251 51.1 52.6 4269 13000 12891 51.1 45 0.1 768 76.5 42.6 36 53.2 44 176 42.2 42.6 42.2 42.6 42.2 42.6 42.2 4					50.5	52.6	4255	3500	3481	51.2	50	0.6	277	74.6			_
739 1757			3224	3242	50.5	52.6	4256	3497	3478	51.3	50	0.8	274				
738 1758 9140 9159 50.1 45 4258 9597 9576 51 40.9 0.9 458 75.3 41.3 36 52.7 740 1760 9140 9159 50.1 45 4260 9560 9540 51.6 42.9 1.5 421 75.2 41.3 36 52.7 741 1761 7728 7746 51.7 52.6 4261 8189 8170 50.6 50 1.1 462 75.7 42 36 53.3 742 1762 9140 9159 50.1 45 4262 3559 5959 50.6 42.9 0.5 42.0 75.3 41.4 36 52.7 742 1762 9140 9159 50.1 45 4262 35.55 52.55 50.6 42.9 0.5 42.0 75.3 41.4 36 52.7 742 1763 12235 12253 50.1 52.6 4263 12998 12979 50.1 45 0.0 764 76.5 42.5 36 53.7 744 1764 3225 3244 52.4 55 4264 3503 3444 61.5 50 1 279 74.8 42.3 36 52.2 746 1768 3225 3244 52.4 55 4268 3500 3481 51.2 50 1.3 276 74.7 42 36 53.2 748 1768 3225 3244 52.4 55 4268 3500 3481 51.2 50 1.3 276 74.7 42 36 52.7 749 1769 12233 12251 51.1 52.6 4268 12999 12980 50.6 40 0.6 767 76.4 42.5 36 53.8 749 1769 12233 12251 51.1 52.6 4268 13000 12981 51.1 45 0.1 768 76.5 42.6 36 53.8 750 1770 28396 28414 51.5 44270 28671 28682 52.8 55 13.3 277 78.5 51.3 36 55.1 752 1772 9831 9950 50.2 45 4272 10449 10431 50.9 47.4 0.8 519 75.3 40.8 36 52.7 752 1773 12235 12253 50.1 52.6 4273 12999 12976 50.3 47.4 0.8 519 75.3 40.8 36 52.7 753 1777 12235 12253 50.1 52.6 4273 12999 12976 50.3 47.4 0.8 519 75.3 40.8 36 52.5 753 1777 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.8 519 75.3 40.8 36 52.5 754 1777 1777 28396 28416 52.4 47.6 4272 28671 28662 52.8 55 0.3 27.7 78.5 51.3 36 55.5 755 1777 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.8 519 76.5 42.5 36	737	1757	1	22	54.8	50	4257	204	185	56.6	55	1.8	_			-	
739 759 28179 28200 50.8 40.9 4259 28671 28653 50.2 52.6 6.6 493 79.7 51.7 30 55.7 740 1760 9140 9159 50.1 45 4260 9560 9540 51.6 42.9 1.5 421 75.2 41.3 36 52.8 741 1761 7728 7746 51.7 52.6 4261 8189 8170 50.6 50 1.1 462 75.7 42 36 52.8 742 1762 9140 9159 50.1 45 4262 9559 9539 50.6 42.9 0.5 420 75.3 41.4 36 52.8 743 1763 12235 12235 50.1 52.6 4263 12998 12979 50.1 45 0.7 64 76.5 42.8 36 53.7 744 1764 3225 3244 52.4 55 4264 3503 3484 51.5 50 1 279 74.8 42.3 36 52.7 746 1766 3225 3244 52.4 55 4266 3503 3484 51.5 50 1 279 74.8 42.3 36 52.7 746 1766 3225 3244 52.2 40 4265 15595 15576 50.8 45 1.3 246 75.5 40.6 36 52.7 747 1767 12233 12251 51.1 52.6 4267 12999 12980 50.6 40 0.6 767 76.4 42.5 36 53.7 748 1768 3225 3244 52.4 55 4268 3497 3478 51.3 50 1.1 273 74.7 42.1 36 52.8 750 1770 28395 28414 51.5 45 4270 28671 28651 51.9 51.1 50.9 47.4 0.1 768 76.5 42.6 36 53.7 751 1771 28395 28414 51.5 45 4270 28671 28653 50.2 52.6 1.3 277 78.5 51.3 36 55.1 752 1772 9301 9950 50.2 45 4272 10449 10431 50.9 47.4 0.8 19 75.3 40.8 36 52.8 753 1773 12255 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.2 760 76.4 42.5 36 53.8 756 1776 1753 11543 11562 50.4 40 4275 12861 1236 1237 1236 1237 1235 12253 1235 50.1 52.6 4273 12994 12976 50.3 47.4 0.2 760 76.4 42.5 36 53.8 759 1777 28396 28416 52.4 47.6 4276 28672 28653 53.1 50.0 47.6 0.4 76.5 42.6 36 53.7 759 1777 179 3299 3248 50.6 50 4278 3647 3628 50.6 50.4 42.9 36 53			9140	9159	50.1	45	4258	9597	9576	51							
T40 1760	739	1759	28179	28200	50.8	40.9	4259	28671	28653	50.2	52.6	0.6					
744 1761 7728 7746 51.7 52.6 24261 8189 8170 50.6 850 1.1 462 75.7 42 36 53.3 742 1762 9140 9159 50.1 45 4262 9559 9539 50.6 42.9 0.5 420 75.3 41.3 36 53.3 744 1764 3225 3224 5225 30.1 52.6 4264 3503 3484 51.5 50 11 279 74.8 42.3 36 52.9 745 1766 3225 3244 52.4 55 4264 3503 3484 51.5 50 11 279 74.8 42.3 36 52.9 746 1766 3225 3244 52.4 55 4266 3500 3484 51.5 50 11 279 74.8 42.3 36 52.9 747 1767 12233 12251 51.1 52.6 4267 12999 12980 50.6 40 0.6 767 76.4 42.5 36 53.8 749 1769 12233 12251 51.1 52.6 4267 12999 12980 50.6 40 0.6 767 76.4 42.5 36 53.8 749 1769 12233 12251 51.1 52.6 4268 13000 12881 51.1 45 0.1 768 765. 42.6 36.5 750 1770 28395 28414 51.5 45 4270 28671 28635 50.2 52.6 427 28637 28671 28635 52.8 55.1 32.7 78.5 51.3 36 55.7 751 1771 28395 28414 51.5 44.9 4271 28637 28652 52.8 55.1 13 277 78.5 51.3 36 55.5 752 1772 9931 9950 50.2 45 4271 28671 28652 52.8 55.1 13 277 78.5 51.3 36 55.5 756 1776 1343 11562 50.4 40 40275 12258 1238 53.1 50.1 19 287 50.4 42.5 36 53.8 755 1777 28396 28416 52.4 47.6 4277 28671 28652 52.8 55.1 50.5 1.9 287 78.5 51.3 36 55.5 756 1777 28396 28416 52.4 47.6 4277 28671 28652 52.8 55.1 50.5 52.6 4268 36.5 756 1776 3229 3248 50.6 50.4	740	1760	9140	9159	50.1	45	4260	9560	9540	51.6	42.9						
742 1762 9140 9159 50.1 45 4262 9559 9539 50.6 42.9 0.5 420 75.3 41.4 36 52.8 743 1763 12235 12253 50.1 52.6 4263 12986 12979 50.1 45 0 764 76.5 42.5 36 53.7 744 1764 3225 3244 52.4 55 4266 3500 3484 51.5 50 1 277 74.8 42.3 36 53.7 746 1766 3225 3244 52.4 55 4266 3500 3481 51.2 50 1.3 276 74.7 42 36 52.7 747 1767 12233 12251 51.1 52.6 4267 3299 12990 50.6 40 0.8 76.7 74.4 42.1 36 53.2 749 1768 3225 3244 52.4 55 4268 3500 3481 51.2 50 1.3 276 74.7 42 36 52.7 747 1767 12233 12251 51.1 52.6 4269 13000 12981 51.1 45 0.1 768 76.5 42.6 36 53.2 749 1769 12233 12251 51.1 52.6 4269 13000 12981 51.1 45 0.1 768 76.5 42.6 36 53.2 576 57.0 770 28395 28414 51.5 45 4271 28671 28663 50.2 52.6 1.3 277 78.5 51.3 36 55.5 756 1771 28395 28414 51.5 45 4271 28671 28663 50.2 52.6 1.3 277 78.5 51.3 36 55.5 752 1773 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.2 760 64 42.5 36 53.4 57.5 47.6 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.7 42.9 42.	741	1761	7728	7746	51.7	52.6	4261	8189	8170	50.6							
744 1764 3225 3245 52.6 4263 12998 12979 50.1 45 0 764 76.5 42.5 36 53.7 744 1764 3225 3244 52.4 55 4264 3503 3484 51.5 50 1 279 74.8 42.3 36 52.9 746 1766 3225 3244 52.4 55 4266 3503 3484 51.5 50 1 279 74.8 42.3 36 52.9 747 1767 12233 12251 51.1 52.6 4266 3503 3481 51.2 50 1.3 247 74.7 42.2 36 52.7 748 1768 3225 3244 52.4 55 4266 3503 3481 51.2 50 1.3 277 76.4 42.5 36 53.3 749 1769 12233 12251 51.1 52.6 4268 13000 12981 51.1 45 0.1 768 76.7 74.7 42.1 36 52.8 749 1769 12233 12251 51.1 52.6 4268 13000 12981 51.1 45 0.1 768 76.5 42.6 36 54.7 750 1770 28395 28414 51.5 45 4270 28671 28653 50.2 52.6 1.3 277 78.5 51.3 36 55.5 752 1772 9931 9950 50.2 45 4272 10449 10431 50.9 47.4 0.8 519 75.3 40.8 36 52.9 753 1773 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.8 519 75.3 40.8 36 52.9 754 1774 3399 3379 51.2 42.9 4274 4350 3631 50.1 50.1 292 75.6 42.8 36 53.6 756 1776 28396 28416 52.4 47.6 4276 28672 28683 51.8 55 0.5 277 78.6 51.6 36 55.7 757 1777 23396 22416 52.4 47.6 4276 28672 28683 51.8 55 0.5 277 78.6 51.6 36 55.7 759 1778 3229 3248 50.6 50 4278 3847 3628 50.6 45 0.4 47.6 4276 28672 28685 50.8 45 0.2 77 78.6 51.6 36 53.7 750 1778 3229 3248 50.6 50 4278 3847 3628 50.6 45 0.4 47.6 4276 28672 28673 51.8 55 0.5 277 78.6 51.4 36 53.7 750 1778 3229 3248 50.6 50 4278 3847 3628 50.6 45 0.4 47.6 47.6 4278 28672 28673 51.8 55 0.5 277 78.6 51.4 36 53.7 750 178 3229 3248 50.6 50 4288 3649 3635 50.8 45 0.5 47.6 47.6 427	742	1762	9140	9159	50.1	45	4262	9559	9539	50.6							
744 1764 3225 3244 52.4 55.5 4264 3503 3484 51.5 50 1 279 74.8 42.3 36 52.7 745 1765 14951 14975 52.2 40 4265 15595 15576 50.8 45 13. 276 74.7 42 36 52.7 746 1766 3225 3244 52.4 55.4 65.4 65.5 63.00 3481 51.2 50 13 276 74.7 42 36 52.7 747 1767 12233 12251 51.1 52.6 4267 12999 12980 50.6 40 0.6 767 76.4 42.5 36 53.8 748 1768 3225 3244 52.4 55.4 42.8 42.8 42.8 42.8 42.8 749 1769 12233 12251 51.1 52.6 4268 3497 3478 51.3 50 1.1 273 74.7 42.1 36 52.8 750 1770 28395 28414 51.5 45 4270 28671 28653 50.2 52.6 1.3 277 78.5 51.3 36 55.1 751 1771 28395 28414 51.5 45 4271 28671 28653 50.2 52.8 53 3277 78.5 51.3 36 55.5 752 1772 9931 9950 50.2 45 4272 2471 247	743	1763	12235	12253	50.1	52.6	4263	12998									
745 1765 14951 14975 52.2 40 4265 15595 15576 50.8 45 1.3 645 75.5 40.6 36 53.2 746 1766 3225 3244 52.4 55 4266 3500 3481 51.2 50 1.3 276 74.7 42 36 52.7 747 1767 12233 12251 51.1 52.6 4267 12999 12996 50.6 40 0.8 76.7 76.4 42.5 36 53.8 748 1768 3225 3244 52.4 55 4268 3497 3476 51.3 50 1.1 273 74.7 42.1 36 52.8 750 1770 28385 28414 51.5 45 4270 28671 28665 50.2 52.6 1.3 277 78.5 51.3 36 55.5 751 1771 28395 28414 51.5 45 4271 28671 28665 50.2 52.6 1.3 277 78.5 51.3 36 55.5 752 1772 9931 9950 50.2 45 4272 10449 10431 50.9 47.4 0.8 519 75.3 40.8 36 52.7 753 1773 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.2 76.4 42.5 36 53.4 756 1776 28396 28416 52.4 47.6 4276 28672 28652 52.8 55 1.3 277 78.5 51.3 36 55.5 756 1776 28396 28416 52.4 47.6 4277 28673 50.3 42.9 0.2 716 76.1 41.9 36 53.5 756 1777 28396 28416 52.4 47.6 4277 28673 28682 52.8 55 0.5 27.7 78.6 51.4 36 53.4 758 1778 3229 3248 50.6 50 4278 3647 3628 50.6 45 0.4 47.6 4277 28673 28682 52.8 55 0.4 27.6 78.6 51.4 36 55.5 759 1779 12235 12253 50.1 52.6 4278 12992 12974 51.2 50.6 41.1 75.3 41.4 36 53.7 759 1779 12235 12253 50.1 52.6 4278 3680 3631 53.1 50 14.4 418 75.3 41.4 36 53.7 750 1780 3229 3248 50.6 50.4 2480 3646 3625 52.4 50.6 41.1 75.8 41.5 36 53.7 750 1780 3229 3248 50.6 50.4 2480 3646 3625 52.4 40.9 14.4 418 75.3 41.4 36 53.7 760 1780 3229 3248 50.6 50.4 4280 3646 3631 53.1 50 1.4 428 75.5 44.8 36 53.7 761 1781 3228 3248 52.4 47.6 4287 3680 3631 53.1 50	744	1764	3225	3244	52.4	55	4264										
746 1766 3225 3244 55.4 55 4266 3500 3481 51.2 50 1.3 276 74.7 42 38 52.7 747 1767 12233 12251 51.1 52.6 4267 12999 12980 50.6 40 0.6 767 76.4 42.5 38 53.8 748 1768 3225 3244 52.4 55.4 55 4288 3497 3478 51.3 50 1.1 273 74.7 42.1 36 52.8 749 1769 12233 12251 51.1 52.6 4268 13000 12981 51.1 45 0.1 788 76.5 42.6 36 5.5 5.7 750 1770 28395 28414 51.5 45 4270 28671 28663 50.2 52.6 1.3 277 78.5 51.3 36 55.5 752 1772 28391 3950 50.2 45 4272 10449 10431 50.9 47.4 0.8 519 75.3 40.8 36 52.9 753 1773 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.8 519 75.3 40.8 36 52.9 753 1773 12235 12253 50.1 52.6 4273 12994 12976 50.3 47.4 0.2 760 76.4 42.5 38 53.4 755 1775 1176 1177 28396 28416 51.5 40.4 4275 12258 12238 50.3 42.9 0.2 776 76.1 42.5 38 53.4 755 1775 11543 11562 50.4 40 4275 12258 12238 50.3 42.9 0.2 776 76.1 41.9 36 53.5 759 1777 28396 28416 52.4 47.6 4276 28672 28653 51.8 55 0.5 277 78.6 51.6 38 55.7 751 1777 28396 28416 52.4 47.6 4276 28672 28653 51.8 55 0.5 277 78.6 51.6 38 55.7 751 1777 28396 28416 52.4 47.6 4276 28672 28653 51.8 55 0.5 277 78.6 51.6 38 55.7 751 1777 28396 28416 52.4 47.6 4276 28672 28653 51.8 55 0.5 277 78.6 51.6 38 55.7 751 1777 28396 28416 52.4 47.6 4276 28672 28653 51.8 55 0.4 276 78.6 42.6 36 53.7 760 1780 3229 3248 50.6 50 4278 3647 3628 50.6 45 0.4 479 75.3 41.5 36 53.7 761 1781 3223 3248 50.6 50 4288 3648 3625 52.8 55 0.4 276 78.6 51.4 36 55.7 760 1780 3229 3248 50.6 50 4280 3646 3625 52.8 55 0.4 276 78.6 51.6 36 53.7 761 1781 3228 3248 50.6 50 4288 30449 10428 51.9 40.9 11.4 418 75.3 41.4 36 52.8 761 1781 3228 3248 50.6 50 4288 30449 50.8 50.8 45 0.0 11.1 423 75.4 41.6 36 53.7 761 1787 2429 2447 50.2 42.9 4284 1622 1602 51.6 67.6 14. 10 57.5 40.8 36 53.7 761 1787 2429 2447 50.2 42.9 4284 1622 1602 51.6 67.6 14. 10 57.5 40.8 36 53.7 761 1787 2429 2447 50.2 42.9 4284 1622 1602 51.6 67.6 14. 10 57.5 40.8 36 53.7 761 1787 2429 2447 50.2 47.4 4281 3860 3631 55.1 50.0 43.6 67 75.8 41.4 36 53.7 771 1799 943 961 50.3 47.4 4291 1483 1444 1466 51.3 45.5 1.5 50.0 486 399 13039 13039 13039 13039 1	745	1765	14951	14975	52.2	40	4265	15595									
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770 1790 942 961 52.8 50 4290 1493 1496 53.1 52.6 0.4 54.3 76.9 44.4 36 54.7 771 1791 943 961 50.3 47.4 4291 1493 1494 51.3 45 1 541 76.8 44.2 36 53.9 772 1792 28867 28866 53.2 50 4292 29358 2339 52.8 50 0.3 492 76.9 44.9 36 53.9 773 1783 943 961 50.3 47.4 4293 29358 29339 52.8 50 0.3 492 76.9 44.9 36 53.9 774 1794 28866 28866 55.4 52.4 4294 29301 29282 55.3 55 0.2 436 77 45.4 36 53.9 775 1795 12352 12375 52.9 41.7 4295 12997 12977 51.8 42.9 1.1 646 76.4 42.9 36 54.1 776 1796 28867 28887 53.7 47.6 4296 29358 29339 52.8 50 0.9 492 76.9 44.9 36 53.2 778 1798 6098 6118 50.3 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.4 779 1799 28868 28888 51.4 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.4 780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1																	
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773 1793 943 961 50.3 47.4 4293 1483 1465 50.5 47.4 0.3 541 76.8 44.2 36 53.9 774 1794 28866 2886 55.4 52.4 4294 28301 29282 55.3 55 0.2 436 77 45.4 36 55.6 775 1795 12352 12352 12375 52.9 41.7 4295 12997 12977 51.8 42.9 1.1 646 76.4 42.9 36 54.1 776 1796 28867 28887 53.7 47.6 4296 29358 29339 52.8 50 0.9 492 76.9 44.9 36 54.8 777 1797 3986 3917 50.7 40.9 4297 4608 4590 51.5 52.6 0.9 713 75.5 40.4 36 53.2 778 1798 6098 6118 50.3 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.2 779 1799 28868 28888 51.4 42.9 4298 6486 6467 50.8 45 0.5 1.4 491 76.9 44.8 36 54.3 780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1																36	
774 1794 28866 28886 55.4 4294 229301 22922 55.3 55 0.2 436 77 45.4 36 55.6 775 1795 12332 12375 52.9 41.7 4295 12997 12977 51.8 42.9 1.1 646 76.4 42.9 36 54.1 776 1796 28867 28887 53.7 47.6 4296 29358 29339 52.8 50 0.9 492 76.9 44.9 36 54.8 777 1797 3896 3917 50.7 40.9 4297 4608 4590 51.5 52.6 0.9 713 75.5 40.4 36 53.2 778 1798 6938 6118 50.3 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.4 779 1799 28868 28888 51.4 42.9 4299 29358 29339 52.8 50 1.4 419 76.9 44.8 36 54.3 780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1 781 1800 2220 2220 5230 54.4 44.000 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1 781 1801 2220 2220 5230 54.4 44.000 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1			_												44.9	36	54.8
775 1795 12382 12375 52.9 41.7 4295 12997 12997 51.8 42.9 1.1 64.5 76.4 42.9 36 54.1 776 1796 28867 28887 53.7 47.6 4298 29358 29339 52.8 50 0.9 492 76.9 44.9 36 54.8 777 1797 3896 3917 50.7 40.9 4297 4608 4590 51.5 52.6 0.9 713 75.5 40.4 36 53.2 778 1798 6098 6118 50.3 42.9 4298 6486 6467 50.8 45 0.5 388 74.6 40.0 36 52.4 790 1890 8220 8240 54 42.9 4299 29358 29339 52.8 50 1.4 491 76.9 44.8 36 54.3 780 1890 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1 781 1890 2220 2220 5220 54.4 44.9 44.9 44.8 64.6 46.6																36	53.9
776 1796 28867 28887 53.7 47.6 4296 22958 22939 52.8 50 0.9 442 76.9 44.9 36 54.1 777 1797 3896 3917 50.7 40.9 4297 4608 4580 51.5 52.6 0.9 713 75.5 40.4 36 53.2 778 1798 6098 6118 50.3 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.2 779 1799 28868 28888 51.4 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.4 780 1890 8220 8240 54 47.6 4300 8831 8913 55.5 52.6 1.4 712 75.4 40 36 54.1 781 1800 8220 8240 54 47.6 4300 8831 8913 55.5 52.6 1.4 712 75.4 40 36 54.1																	55.6
777 1797 3896 3917 50.7 40.9 4297 4608 4590 51.5 52.6 0.9 71.3 75.5 40.4 36 53.2 778 1798 6098 6118 50.3 42.9 4298 6486 6467 50.8 45 0.5 389 74.6 40.1 36 52.4 779 1799 28888 2888 51.4 42.9 4299 29358 29339 52.8 50 1.4 491 76.9 44.8 36 54.3 780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1															42.9	36	54.1
778 1798 6098 6118 50.3 42.9 4299 6486 6467 50.8 45 0.5 389 74.6 40.1 36 53.2 779 1799 28868 2888 51.4 42.9 4299 29358 29339 52.8 50 1.4 491 76.9 44.8 36 54.3 780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1 781 1801 2220 2230 54.2 45 4400 8931 8933 55.5 52.6 1.4 712 75.4 40 36 54.1															44.9	36	54.8
779 1799 28868 2888 51.4 42.9 4299 29958 29358 2939 52.8 50 1.4 491 76.9 44.8 36 54.1 780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1 781 1801 2220 2220 51.2 45 440 20 20 20 20 20 20 20 20 20 20 20 20 20														75.5	40.4	36	53.2
780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1												0.5	389	74.6	40.1	36	52.4
780 1800 8220 8240 54 47.6 4300 8931 8913 55.5 52.6 1.4 712 75.4 40 36 54.1													491	76.9	44.8	36	54.3
781 1801 2220 2239 51.3 45 4301 2672 2653 51.6 50 0.4 453 77 45.3 36 54.4											52.6	1.4	712	75.4	40	36	
	/81	1801	2220	2239	51.3	45	4301	2672	2653	51.6	50	0.4	453	77	45.3	36	54.4

782	1802	12040	12057	50.6	50	4302	12493	12476	50.7	50	0.1	454	76.3	43,6	36	53.7
783	1803	942	960	52.1	52.6	4303	1483	1464	51.3	45	0.8	542	76.8	44,3	36	54.3
784	1804	28868	28889	52	40.9	4304	29358	29339	52.8	50	0.8	491	76.9	44.8	36	54.5
785	1805	942	960	52.1	52.6	4305	1483	1465	50.5	47.4	1.6	542	76.8	44.3	36	54
786	1806	12040	12057	50.6	50	4306	12724	12705	52.4	55	1.8	685	76.6	43.1	36	53.9
787	1807	942	960	52.1	52.6	4307	1484	1466	53.1	52.6	1	543	76.9	44.4	36	54.5
788	1808	11545	11563	50.8	47.4	4308	12253	12235	50.1	52.6	0.7	709	76.2	42.2	36	53.5
789	1809	98	118	50.6	42.9	4309	269	251	51.1	52.6	0.5	172	75	46.5	36	52.8
790	1810	12373	12391	50.8	47.4	4310	12911	12892	50.5	50	0.3	539	76.1	42.5	36	53.5
791	1811	16366	16384	50.3			16781	16761	51.3	47.6	0.9	416	75.1	41.1	_36	52.8
792		9929	9946	50	50	4312	10183	10165	51.7	47.4	1.6	255	75.3	43.9	36	52.8
793		12236	12256	51.2		4313	13000	12981	51.1	45	0.1	765	76.4	42.5	36	53.9
794	1814	3231	3252	52.7		4314	3650	3631	53.1	50	0.4	420	75.4	41.7	36	53.7
795		11541	11560	50.1		4315	11727	11708	50.4	45	0.3	187	73	40.6	36	51.2
796		3232	3252	51.1		4316	3494	3473	50.4	40.9	0.7	263	74.3	41.4	36	52.3
797	1817	7725	7743	50.8		4317	8054	8035	50.4	50	0.4	330	75	41.8	36	52.7
798		28968	28988	50.9			29358	29339	52.8	50	2	391	76.4	44.5	36	53.9
799		11545	11563	50.8		4319	12257	12237	51.3	47.6	0.5	713	76.2	42.1	36	53.7
800		24417	24436	52.6			25080	25062	53.5	52.6	0.9	664	75.8	41.3	36	53.9
801	1821	28968	28989	51.5		4321	29358	29339	52.8	50	1.3	391	76.4	44.5	36	54.1
		3789	3808	53.5		4322	4318	4294	54.4	40	0.9	530	75.5	41.1	36	54.1
803		3232	3252	51.1		4323	3646	3625	52	40.9	0.9	415	75.3	41.4	36	53.1
804		3232	3252	51.1		4324	3647	3628	50.6	45	0.5	416	75.3	41.6	36	53
805		28971	28993	51.9		4325	29306	29288	53.5	52.6	1.6	336	76.2	44.6	36	54
806		24179	24200	53.3		4326	24818	24797	51.6	40.9	1.6	640	75.8	41.2	36	53.6
807	1827	3231	3251	52	47.6		3650	3631	53.1	50	1.1	420	75.4	41.7	36	53.5
808		9930	9950	52.6			10449	10425	54.6	40	2	520	75.4	41.7	36	53.7
		8866	8885	51.1		4329	9254	9236	50.6	47.4	0.5	389	75	41.1	36	52.8
	1830	2522	2541	51.4		4330	2672	2653	51.6	50	0.3	151	75.3	48.3	36	53.2
811	1831	11541	11561	50.9		4331	12258	12238	50.3	42.9	0.6	718	76.2	41.9	36	53.5
812	1832	3232	3251	50.3		4332	3646	3625	52	40.9	1.7	415	75.3	41.4	36	52.9
	1833	3232	3251	50.3		4333	3647	3628	50.6	45	0.3	416	75.3	41.6	36	52.9
	1834	23843	23863	50.3		4334	24527	24508	50.5	45	0.3	685	76.3	41.8	36	53.4
815	1835	21210	21228	53.2	52.6		21317	21293	53.2	40	0.2	108	71.1	41.8	$\overline{}$	
		3229	3249	51.4		4336	3650	3631	53.1	50	1.7	422	75.4	41.7	36 36	50.8 53.3
817	1837	3230	3249	50.1		4337	3494	3473	50.4	40.9	0.3	265	74.2	41.1	36	52.1
		2371	2389	50.3		4338	2997	2976	51.4	40.9	1.1	627	76.7	43.5	36	53.9
		29186	29206	51.3		4339	29298	29280	51.4	52.6	0.1	113	72.8	43.5	36	51.5
		9929	9946	50		4340	10455	10435	50.5	42.9	0.1	527	75.3	40.6	36	52.8
821	1841	9351	9370	51.2		4341	9989	9968	51	40.9	0.4	639	75.4	40.6	36	53.2
		25348	25365	50.4		4342	25772	25753	51.9	50	1.5	425	74.9	40.4	36	52.6
		1402	1422	50.2		4343	2103	2082	52	45.5	1.8	702	76.7	43.3	36	53.8
824	1844	9929	9946	50		4344	10608	10589	51	50	0.9	680	75.8	41.2	36	53.2
	1845	9934	9953	50.7		4345	10608	10589	51	50	0.3	675	75.8	41.2	36	53.4
826	1846	13176	13196	51.4		4346	13544	13525	52.6	55	1.2	369	77.1	46.3	36	54.5
827	1847	7725	7743	50.8		4347	8189	8170	50.6	50	0.2	465	75.6	41.9	36	
828	1848	7725	7743	50.8		4348	8190	8172	50.8	47.4	0.2	466	75.6	41.8		53.2 53.1
829	1849	18074	18093	50.3		4349	18662	18641		40.9					36	
830	1850	18074	18093	50.3		4350	18632	18611	50.4	40.9	0.1	589 559	76.2	42.6	36	53.6
831	1851	29200	29222	53.2		4351	29306	29288	53.5	52.6			76.2	42.8	36	53.5
832	1852	25348	25366	51.2		4352	25545	25526	51.7	45	0.3	107	73.1	47.7	36	52.3
002	1002	20040	20300	01.2	47.4	4002	20545	20026	51.7	45	0.5	198	74.1	42.9	36	52.3

833	1853	25348	25366	51.2	47.4	4353	25545	25525	52.3	42.9	1.1	198	74.1	42.9	36	52.3
834	1854	29200	29223	53.7	41.7	4354	29306	29288	53.5	52.6	0.2	107	73.1	47.7	36	52.3
835	1855	25347	25366	52.7	50	4355	25545	25521	54.5	40	1.8	199	74.2	43.2	36	52.9
836	1856	3792	3811	54	55	4356	4447	4425	53	43.5	1	656	75.5	40.5	36	53.9
837	1857	29200	29224	54.2	40	4357	29306	29288	53.5	52.6	0.7	107	73.1	47.7	36	52.3
838	1858	985	1004	51.1	50	4358	1483	1465	50.5	47.4	0.6	499	76.4	43.5	36	53.7
839	1859	2427	2445	52.1	52.6	4359	3189	3168	51	45.5	1.1	763	76.7	43.1	36	54.1
840	1860	13701	13725	53.6	40	4360	14084	14060	53.6	40	0.1	384	74.6	40.1	36	53.4
841	1861	985	1004	51.1	50	4361	1483	1464	51.3	45	0:2	499	76.4	43.5	36	53.9
842	1862	8794	8813	51.6	45	4362	9559	9539	50.6	42.9	1	766	75.9	41.3	_ 37	53.4
843	1863	3789	3806	50	50	4363	4435	4417	50.5	52.6	0.4	647	75.5	40.5	37	52.9
844	1864	13177	13197	50.3	42.9	4364	13314	13297	51	50	0.6	138	72.8	43.5	37	51.2
845	1865	3791	3808	50	50	4365	4435	4417	50.5	52.6	0.4	645	75.4	40.5	37	52.9
846	1866	9139	9159	52.5	47.6	4366	9364	9346	53.9	52.6	1.4	226	74.9	43.8	37	53.3
847	1867	3226	3245	51.7	55	4367	3494	3473	50.4	40.9	1.3	269	74.5	41.6	37	52.4
848	1868	13040	13059	50.9	50	4368	13314	13297	51	50	0.1	275	75.6	44.4	37	53.3
849	1869	2522	2541	51.4	45	4369	2891	2873	50.8	47.4	0.6	370	76	43.8	37	53.6
850	1870	8865	8884	50.4	45	4370	9245	9226	50	45	0.4	381	74.9	40.9	37	52.6
851	1871	3787	3804	50	50	4371	4434	4416	51.5	52.6	1.4	648	75.4	40.3	37	52.9
852	1872	3226	3245	51.7	55	4372	3646	3625	52	40.9	0.3	421	75.3	41.6	37	53.4
853	1873	3226	3245	51.7	55	4373	3647	3628	50.6	45	1.1	422	75.4	41.7	37	53.1
854	1874	3226	3245	51.7	55	4374	3650	3631	53.1	50	1.4	425	75.5	41.9	37	53.5
855	1875	2387	2405	51.6	52.6	4375	2747	2727	50	42.9	1.6	361	76.9	46	37	53.9
856	1876	18074	18093	50.3	45	4376	18229	18209	50.1	42.9	0.2	156	73.2	42.9	37	51.4
857	1877	13701	13725	53.6	40	4377	14059	14040	52.8	50	0.8	359	74.5	40.1	37	53.1
858	1878	3787	3804	50		4378	4435	4417	50.5	52.6	0.4	649	75.4	40.4	37	52.9
859	1879	13040	13059	. 50.9	50	4379	13323	13304	51.1	45	0.2	284	75.7	44.4	37	53.4
860	1880	3789	3806	50		4380	4434	4416	51.5	52.6	1.4	646	75.4	40.4	37	52.9
861	1881	15506	15527	50.8	40.9	4381	16214	16196	51.8	52.6	1	709	75.5	40.3	37	53.2
862	1882	12234	12252	50.6	47.4	4382	12412	12392	50	42.9	0.6	179	73.1	41.3	37	51.3
863	1883	12234	12252	50.6	47.4	4383	12739	12718	51	40.9	0.4	506	75.8	42.1	37	53.4
864	1884	18074	18094	51.1	42.9	4384	18229	18209	50.1	42.9	0.9	156	73.2	42.9	37	51.4
865	1885	18075	18095	50.6	47.6	4385	18223	18206	51.8	50	1.2	149	73.3	43.6	37	51.6
866	1886	13040	13059	50.9	50	4386	13326	13306	50.7	42.9	0.2	287	75.7	44.3	37	53.3
867	1887	18080	18098	51.2	52.6	4387	18233	18215	51.3	52.6	0.1	154	73.9	44.8	37	52.2
868	1888	18080	18098	51.2	52.6	4388	18233	18214	52	50	0.9	154	73.9	44.8	37	52.2
869	1889	18080	18098	51.2	52.6	4389	18231	18210	52.2	45.5	1	152	73.5	44.1	37	51.9
870	1890	18080	18098	51.2	52.6	4390	18223	18206	51.8	50	0.6	144	73.2	43.8	37	51.7
871	1891	18077	18098	52.9		4391	18220	18202	54.8	52.6	1.9	144	73.2	43.8	37	52.2
· 872	1892	18076	18098	54.4		4392	18443	18424	55.9	55	1.5	368	75.8	43.2	37	54.5
873	1893	3792	3810	52.9		4393	4436	4417	52.2	50	0.6	645	75.4	40.5	37	53.6
874	1894	3055	3074	51.1		4394	3647	3628	50.6	45	0.5	593	76.3	42.7	37	53.6
875	1895	3055	3074	51.1		4395	3646	3625	52	40.9	0.9	593	76.2	42.7	37	53.7
876	1896	15506	15527	50.8	40.9		15645	15625	51.1	42.9	0.4	140	71.8	40.7	37	50.6
877	1897	18081	18099	51.2		4397	18229	18209	50.1	42.9	1.1	149	73.3	43.6	37	51.4
878	1898	13039	13058	51.8		4398	13155	13138	50.4	50	1.4	117	73.4	43.6	37	51.6
879	1899	3055	3075	51.8	47.6		3650	3631	53.1	50	1.3	596	76.3	42.8	37	54.1
880	1900	18080	18099	53		4400	18223	18205	53.3	52.6	0.4	144	73.2	43.8	37	52.2
881	1901	27361	27380	52.4		4401	27579	27558	51.1	40.9	1.3	219	75.4	45.8	37	
882	1902	3221	3239	51.5	52.6		3503	3484	51.5	50	0	283	74.8	45.2	37	53.2
883	1903	3221	3239	51.5	52.6		3504	3485	50.4	45	1.1	284	74.7	41.9	37	52.9
				- 1101	-3.0		3004	0400	50.4	40	1.11	204	14.1	41.9	3/	52.5

884	1904	18077	18099	54.4	47.8	4404	18220	18201	56.1	55	1.7	144	73.2	43.8	37	52.6
885	1905	3055	3075	51.8	47.6	4405	3647	3628	50.6	45	1.2	593	76.3	42.7	37	53.7
886	1906	18581	18599	51.4	47.4	4406	18697	18679	51.9	52.6	0.4	117	71	41	37	50.2
887	1907	18616	18636	51.4	47.6	4407	19216	19195	50.2	40.9	1.1	601	75.6	41.1	37	53.1
888	1908	3219	3238	50.7	50	4408	3503	3484	51.5	50	0.8	285	74.8	42.1	37	52.7
889	1909	18696	18715	51.7	. 50	4409	19216	19195	50.2	40.9	1.5	521	75.5	41.3	37	53
890	1910	3219	3238	50.7	50		3504	3485	50.4	45	0.3	286	74.7	42	37	52.5
891	1911	27366	27384	52.2	52.6	4411	27573	27552	52.3	40.9	0.1	208	74.6	43.8	37	53
892		27366	27384	52.2		4412	27567	27547	51.1	42.9	1	202	74.6	44.1	37	52.7
	1913	3055	3075	51.8		4413	3646	3625	52	40.9	0.2	592	76.2	42.6	_37	54.7
894		18704	18724	50.8		4414	19216	19195	50.2	40.9	0.5	513	75.5	41.1	37	53
895		16874	16893	52.1		4415	17056	17035	51.8	45.5	0.4	183	74.4	44.3	37	52.7
	1916	12234	12252	50.6		4416	12739	12719	50.3	42.9	0.4	506	75.8	42.1	37	53.3
897	1917	7728	7746	51.7	52.6		8054	8035	50.4	50	1.2	327	75.0	41.9	37	
898		15506	15527	50.8	. 40.9		15647	15628	51	45	0.3	142	71.9	40.8		52.7
899		985	1004	51.1		4419	1773	1755	51.7	52.6	0.6	789	76.7	43.1	37	50.7 54.1
900		3217	3236	51.1	50		3503	3484	51.5	50	0.4	287	74.8			
901	1921	3791	3808	50		4421	4434	4416	51.5	52.6	1.4	644		42.2	37	52.8
902	1922	19794	19813	50	50		19923	19904	50.1	52.6 50	0.1		75.4	40.4	37	52.9
903		13039	13058	51.8	50		13178	13157	50.1			130	72.7	43.8	37	51
904		13033	13051	52.1	52.6		13175	13138	50.4	40.9 50	1.5	140	73.8	45.7	37	51.9
905		12233	12251	51.1	52.6		12739	12719	50.4		1.7	123	73.7	47.2	37	51.8
906	1926	19795	19814	50.4	45	4425	19923	19903		42.9	0.9	507	75.9	42.2	37	53.3
907	1927	13177	13197	50.4		4426	13946		50.9	47.6	0.4	129	72.5	43.4	37	51
908	1928	3799	3820	52.9	45.5			13929	51.5	50	1.2	770	75.9	41	37	53.3
909	1928	8867					4318	4294	54.4	40	1.5	520	75.4	41	37	53.7
910	1930	1472	8887 1491	52.3	47.6		9364	9346	53.9	52.6	1.6	498	75.8	42.2	37	53.9
		12233		51.2	45	4430	2152	2133	50.7	45	0.5	681	76.5	42.9	37	53.8
911	1931 1932	3055	12251	51.1		4431	12739	12718	51	40.9	0.2	507	75.9	42.2	37	53.5
912	1932	12726	3076 12746	52.4	45.5	4432	3650	3631	53.1	50	0.7	596	76.3	42.8	37	54.2
	1933	8867		51.3		4433	13325	13305	50.5	47.6	0.7	600	76.7	43.7	37	53.9
			8887	52.3		4434	9316	9296	50.8	42.9	1.5	450	75.4	41.6	37	53.1
915	1935	8867 8867	8887	52.3		4435	9314	9295	51.1	50	1.2	448	75.5	41.7	37	53.3
917	1936		8887	52.3		4436	9313	9294	50.4	50	1.9	447	75.5	41.6	37	53
	1937 1938	3055	3076	52.4	45.5	4437	3647	3628	50.6	45	1.8	593	76.3	42.7	37	53.7
		13176	13196	51.4		4438	13312	13294	51	52.6	0.4	137	72.9	43.8	37	51.5
919		12726	12746	51.3			13155	13138	50.4	50	0.9	430	76.4	44	37	53.7
920 921	1940 1941	13701	13724	53.1	41.7	4440	14058	14040	51.4	52.6	1.7	358	74.5	40.2	37	52.7
921		8372	8390	50.7	47.4		9101	9081	50.5	47.6	0.2	730	75.5	40.3	37	53.1
	1942	3055	3076	52.4		4442	3646	3625	52	40.9	0.4	592	76.2	42.6	37	54.1
923	1943	887	905	50.1		4443	1493	1474	50.8	45	0.7	607	77.1	44.6	37	54.1
924	1944	1046	1063	50.3	50	4444	1697	1677	51	42.9	0.7	652	76.9	43.9	37	54
925	1945	27378	27397	50.5	45		27675	27656	50	40	0.5	298	74.1	40.3	37	52
926	1946	27378	27397	50.5	_ 45	4446	27674	27654	51.9	42.9	1.4	297	74.2	40.4	37	52.2
927	1947	2671	2692	52.1		4447	3056	3037	52.1	55	0	386	74.8	40.7	37	53.1
928	1948	1046	1063	50.3			1697	1678	50.3	45	0.1	652	76.9	43.9	37	54
929	1949	2387	2405	51.6		4449	2672	2654	50.9	52.6	0.8	286	77	47.6	37	54.3
930	1950	3792	3810	52.9		4450	4565	4542	53.9	41.7	1	774	75.6	40.3	37	53.8
931	1951	15506	15527	50.8		4451	15647	15629	50.3	47.4	0.5	142	71.9	40.8	37	50.5
	1952	8794	8813	51.6		4452	9560	9540	51.6	42.9	0	767	75.9	41.2	37	53.7
933	1953	19801	19819	53.2			19909	19885	52.5	40	0.7	109	71.4	43.1	37	50.8
934	1954	19988	20006	50.4	47.4	4454	20615	20597	50.6	47.4	0.2	628	75.3	40.1	37	52.9

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L		1955	19991	20009			4455	20616	20597	52.3	45	0.5	626	75.3	40.	37	53.5
L	936		16875	16895	51.6	47.6	4456	17060	17041	51.1	50				44.6		
L	937		16875	16895	51.6	47.6	4457	17059	17039	50.6	47.6	1			44.3		
L	938		16875	16895	51.6	47.6	4458	17056	17035	51.8	45.5	0.2			4/		
L	939	1959	27442	27461	51.5	40	4459	27541	27521	51.7	47.6	0.1	100		44		
L	940		16875	16896	52.2	45.5	4460	17060	17041	51.1	50	1.1	186	74.6	44.6		52.6
L	941	1961	23841	23859	50.5	52.6	4461	24527	24507	51	42.9	0.5	687	76.1	41.9		53.5
L	942	1	23841	23859	50.5	52.6		24093	24075	50.9	52.6	0.4	253	76	45.8		53.5
L	943		16875	16896	52.2	45.5	1	17059	17039	50.6	47.6	1.6	185	74.4	44.3		52.4
L	944		16875	16896	52.2	45.5		17056	17035	51.8	45.5	0.5	182	74.2	44		52.6
1	945		23843	23863	50.3	42.9	4465	24093	24075	50.9	52.6	0.5	251	75.8	45.4	37	53.3
L	946		16875	16896	52.2	45.5		17041	17023	53.5	52.6	1.3	167	73.8	43.7	37	52.4
L	947	1967	28187	28205	53.1		4467	28673	28654	53.5	55	0.5	487	80	52.4	37	57
L	948		28190	28208	51.7		4468	28672	28654	50.6	52.6	1.2	483	79.9	52.2	37	56.2
⊢	949		24030	24047	50.7	50		24527	24508	50.5	45	0.2	498	75.4	41.2	37	53
H	950		24031	24050	56.5	55		24816	24792	54.7	40	1.8	786	76.3	42	37	54.9
Ļ	951	1	7880	7900	50.3	42.9	4471	8049	8032	50.4	50	0	170	72.8	41.2	37	51.2
-	952		24096	24119	54.4	41.7	4472	24815	24791	54.5	40	0.1	720	75.8	41	37	54.5
-	953 954		17790	17811	51.6	40.9	4473	18233	18214	52	50	0.4	444	75.1	40.8	37	53.2
\vdash		1974	24174	24194	50.9	42.9		24938	24921	50.4	50	0.5	765	75.8	40.9	37	53.3
\vdash	955 956		16875	16896	52.2			17039	17022	51.4	50	0.8	165	73.7	43.6	37	52.1
\vdash	957	1976	24174 24179	24195	52.5			. 24936	24919	51.8	50	0.7	763	75.8	41	37	53.7
\vdash	958	-	16875	24198	51	45	4477	24938	24921	50.4	50	0.6	760	75.8	40.9	37	53.3
-	959	1978	24180	16896 24199	52.2		4478	17038	17021	50.7	50	1.6	164	73.8	43.9	37	52
\vdash	960	1980	2823		50.3		4479	24936	24919	51.8	50	1.5	757	75.8	41	37	53.2
\vdash	961	1981	10142	2844 10163	50.4 51.3	45.5 40.9	4480	3186	3165	50.4	40.9	0	364	75.5	42.6	37	53.1
\vdash	962	1982	1046	1063	50.3		4481 4482	10608	10589	51	50	0.3	467	74.9	40	. 37	52.8
\vdash	963	1983	17388	17408	50.3		4482	1483	1464	51.3	45	0.9	438	76.2	43.6	37	53.6
\vdash	964	1984	24179	24200	53.3	_		17501	17481	51.2	42.9	0.6	114	70.5	40.4	37	49.7
\vdash	965	1985	24379	24398	55		4484 4485	24740	24717	52.5	41.7	0.8	562	76	42.2	37	54
\vdash	966	1986	24379	24398	55		4485	25088 25087	25070	54.5	52.6	0.5	710	75.9	41.3	37	54.6
\vdash	967	1987	16874	16893	52.1		4487	17059	25069 17039	53.7	52.6	1.3	709	75.9	41.3	37	54.3
\vdash	968	1988	24380	24399	55		4488	25088		50.6	47.6	1.5	186	74.6	44.6	37	52.5
-	969	1989	28522	28542	50.2		4489	28671	25070 28653	54.5	52.6	0.5	709	75.9	41.3	37	54.6
\vdash	970	1990	16874	16893	52.1		4490	17060	17041	50.2 51.1	52.6	0	150	76.2	50.7	37	53.5
\vdash	971	1991	24380	24399	55		4491	25087	25069	53.7	50 52.6	1	187	74.7	44.9	37	52.7
r	972	1992	17608	17627	50.2		4492	18239	18220	50.7	45	1.3	708 632	75.9	41.4	37	54.4
\vdash	973	1993	17608	17627	50.2		4493	18238	18219	50.3	45	0.2		75.3	40.2	37	52.8
	974	1994	1046	1063	50.3		4494	1483	1465	50.5	47.4	0.1	631 438	75.3	40.3	37	52.9
	975	1995	17608	17628	50.9	42.9		18239	18220	50.5	45	0.2	632	76.2 75.3	43.6	37	53.6
	976	1996	17608	17628	50.9	42.9		18238	18219	50.3	45	0.5	631	75.3	40.2 40.3	37	52.8
	977	1997	8063	8084	51.4	45.5		8188	8169	50.5	45	0.7	126	72.1	40.3	37 37	52.9
	978	1998	12236	12256	51.2		1498	12739	12718	51	40.9	0.9	504	75.8	42.9		50.7
	979	1999	13176	13196	51.4	47.6		13325	13305	50.5	47.6	0.2	150	73.4	42.1	37	53.5
	980	2000	2371	2389	50.3		4500	2749	2728	50.3	45.5	0.0	379	76.9	45.9	37	51.7
Г	981	2001	9402	9420	51.3		4501	9989	9968	51	40.9	0.4	588	75.4	45.9		
Г	982	2002	9931	9950	50.2		1502	10183	10166	50.9	50	0.4	253	75.2	43.9	37	53.1 52.8
Г	983	2003	2387	2405	51.6	52.6		2997	2976	51.4	40.9	0.7	611	76.6	43.5	37	54.2
	984	2004	3788	3805	50		1504	4435	4417	50.5	52.6	0.4	648	75.4	40.4	37	
	985	2005	26039	26057	52.6	52.6		26650	26630	51.4	42.9	1.2	612	75.3	40.4	37	52.9 53.3
_										24	72.0	1,2)	012	10.0	+0.4	31	03.3

	2006	2371	2389	50.3		4506	3053	3034	50.3	50	0	683	76.7	43.3	37	53.9
987	2007	3	21	53.4	52.6	4507	315	296	51.9	50	1.5	313	76.9	46.6	37	54.5
988	2008	2371	2389	50.3	47.4	4508	3056	3037	52.1	55	1.7	686	76.7	43.4	37	53.9
989	2009	13040	13059	50.9	50	4509	13155	13137	52.1	52.6	1.2	116	73.2	46.6	37	51.6
990	2010	9931	9950	50.2	45	4510	10183	10165	51.7	47.4	1.5	253	75.2	43.9	37	52.8
991	2011	3788	3805	50	50	4511	4434	4416	51.5	52.6	1.4	647	75.4	40.3	37	52.9
992	2012	13176	13196	51.4	47.6	4512	13946	13929	51.5	50	0.2	771	75.9	41.1	37	53.6
993	2013	3772	3792	51.2	42.9	4513	4444	4424	50.6	42.9	0.7	673	75.6	40.7	37	53.2
994	2014	13176	13196	51.4	47.6	4514	13320	13300	51.4	47.6	0	145	73.3	44.1	37	51.9
995	2015	8861	8880	50.2		4515	9245	9226	50	45	0.1	385	74.9	40.8	.37	52.5
996	2016	8868	8889	50.4		4516	9310	9291	51.2	45	0.8	443	75.3	41.3	37	52.9
997	2017	16366	16384	50.3		4517	16774	16752	52.2	43.5	1.9	409	75.1	41.1	37	52.8
998	2018	9934	9953	50.7	50		10183	10166	50.9	50	0.2	250	75.1	44	37	53
	2019	9055	9079	52.8		4519	9342	9323	52.1	50	0.8	288	75.1	42.7	37	53.3
1000		8868	8889	50.4		4520	9249	9231	50.8	47.4	0.6	382	75.1	42.7	37	
1001	2021	16366	16385	52.9		4521	16774	16752	52.2	43.5						52.8
1002		8868	8889	50.4		4522	9249	9230	52.2 51.5		0.6	409	75.1	41.1	37	53.3
1002		9934	9953	50.4		4522	10183			45	1.1	382	75.1	41.4	37	52.8
1003		25772	25793	52.4		4523		10165	51.7	47.4	0.9	250	75.2	: 44	37	53
1004		13039	13057	51.1	52.6		26183 13155	26163 13138	51.7	42.9	0.7	412	74.8	40.3	37	53
		25771	25790	51.1					50.4	50	0.7	117	73.4	47	37	51.6
1000	2020	25769	25786	50.3		4526	26183	26164	51	45	0.1	413	74.8	40.4	37	52.8
					50		26183	26163	51.7	42.9	1.4	415	74.9	40.5	37	52.6
		3794	3812	52.9		4528	4436	4417	52.2	50	0.6	643	75.4	40.4	37	53.6
1009		887	905	50.1	47.4		1480	1462	51.6	47.4	1.5	594	77.1	44.6	37	54.1
		3794	3812	52.9		4530	4434	4416	51.5	52.6	1.4	641	75.4	40.4	37	53.3
		12370	12388	50.1	47.4		12994	12976	50.3	47.4	0.3	625	76.4	42.9	37	53.6
	2032	3797	3815	50.9	47.4		4186	4168	51.8	52.6	0.9	390	75.3	41.8	37	53.1
1013		3795	3813	52.1		4533	4435	4417	50.5	52.6	1.6	641	75.5	40.6	37	53.1
1014		3795	3813	52.1	52.6		4434	4416	51.5	52.6	0.6	640	75.4	40.5	37	53.3
1015		13177	13197	50.3		4535	13323	13304	51.1	45	0.8	147	73.2	43.5	37	51.4
		1046	1064	51.2		4536	1401	1382	50.6	45	0.6	356	75.6	. 43	37	53.2
		16549	16567	54.9	52.6	4537	17057	17035	53	43.5	1.9	509	75.9	42.2	37	54.1
1018	2038	1046	1064	51.2	47.4	4538	1483	1464	51.3	45	0.1	438	76.2	43.6	37	53.8
1019	2039	16551	16568	51.1	50	4539	17056	17035	51.8	45.5	0.7	506	75.9	42.3	37	53.6
		1046	1064	51.2	47.4	4540	1483	1465	50.5	47.4	0.6	438	76.2	43.6	37	53.6
1021	2041	1046	1064	51.2	47.4	4541	1484	1466	53.1	52.6	2	439	76.3	43.7	37	53.9
1022	2042	1046	1064	51.2	47.4	4542	1697	1676	51.7	40.9	0.5	652	76.9	43.9	37	54.2
1023	2043	1046	1064	51.2	47.4	4543	1697	1677	51	42.9	0.2	652	76.9	43.9	37	54.2
1024	2044	16555	16572	50.3		4544	17111	17090	51.1	40.9	0.8	557	76.1	42.5	37	53.5
1025	2045	1046	1064	51.2		4545	1697	1678	50.3	45	0.9	652	76.9	43.9	37	54
1026	2046	1046	1063	50.3		4546	1401	1382	50.6	45	0.2	356	75.6	43	37	53.1
1027	2047	3796	3814	50.8		4547	4435	4417	50.5	52.6	0.3	640	75.4	40.5	37	53.1
1028	2048	12232	12250	51.9		4548	12993	12975	51.4	47.4	0.5	762	76.5	42.5	37	54
1029		12236	12256	51.2		4549	12739	12719	50.3	42.9	0.9	504	75.8	42.5	37	53.2
		28937	28956	52.4		4550	29306	29288	53.5	52.6	1.1	370	76.6	45.1	37	54.4
		12232	12250	51.9	52.6		12996	12977	50.2	40	1.7	765	76.4			
		3234	3254	51.1		4552	3504	3485	50.2	40	0.7	271		42.4	37	53.6
	2053	3234	3254	51.1		4553	3503	3484					74.4	41.3	37	52.3
	2054	9922	9941	51.3		4554	10670	10649	51.5 51.3	50	0.4	270	74.4	41.5	37	52.5
		3792	3810	52.9			4434			40.9	0.1	749	75.8	40.9	37	53.5
		3792	3254	51.1		4555 4550		4416	51.5	52.6	1.4	643	75.4	40.4	37	53.3
		3634	3434	51.1	47.6	4556	3494	3473	50.4	40.9	0.6	261	74.1	41	37	52.1

1037	2057	4255	4276	51.7	45.5	4557	4608	4590	51.5	52.6	0.2	354	74.7	40.7	37	52.8
1038	2058	24562	24580	50.1	52.6		24936	24919		50	1.7	375	75.6		37	
1039	2059	24562	24580	50.1	52.6		24938	24921	50.4	50	0.3		75.5		37	53
1040	2060	24562	24580	50.1	52.6		25182	25164	51.4	47.4	1.3		75.9	41.7	_	53 53.3
1041	12061	24559	24579	52	52.4		24936	24919	51.8	50	0.2	378	75.7	41.7	37	
1042	2062	24559	24579	52		4562	24938	24921	50.4	50	1.6		75.6		37	53.6
1043	2063	1046	1063	50.3	50		1697	1676	51.7	40.9	1.3	652		42.6	37	53.1
1044		24482	24503	51.6		1.000	24815	24792	53.4	41.7	1.8	334	76.9	43.9	37	54
1045		13177	13197	50.3	42.9		13326	13306	50.7	41.7	0.4	150	75.4	42.8	37	53.4
1046		24480	24502	54.2	47.8		24815	24791	54.5	42.9			73.2	43.3	37	51.4
1047		17840	17859	50.8	45		18223	18206	51.8	50	0.3	336	75.6	43.2	37	54.3
1048		24480	24500	53.2		4568	24815	24792	53.4	41.7	1		74.7	40.4	37	52.6
1049		17840	17859	50.8	45		18231	18210			0.2	336	75.6	43.2	37	54
1050		28821	28840	51.8	45		29298		52.2	45.5	1.4	392	74.8	40.6	37	52.7
1051		24418	24440	55	47.8		24517	29279	52.6	55	0.8	478	77	45.2	37	54.6
1052		13701	13722	50.4	40.9		14058	24494 14040	53.2	41.7	1.8	100	70.8	43	37	50.6
	2073	28821	28840	51.8	40.9				51.4	52.6	1	358	74.5	40.2	37	52.4
	2074	17792	17813	51.6	40.9		29358 18233	29339	52.8	50	1	538	77.1	45	37	54.6
	2075	24420	24440	50.8	40.9		25079	18214	52	50	0.4	442	75.1	40.7	37	53.1
	2076	28821	28839	51.1		4576		25061	52.7	52.6	1.9	660	75.7	41.1	37	53.3
1057		3796	3814	50.8		4577	29298	29279	52.6	55	1.5	478	77	45.2	37	54.3
	2078	28821	28839	51.1			4434	4416	51.5	52.6	0.7	639	75.4	40.4	37	53.1
1059		28820	28838	53.7		4578 4579	29358	29339	52.8	50	1.7	538	77.1	45	37	54.4
	2080	24418	24439	52.9	45.5		29298	29279	52.6	55	1.1	479	77.1	45.3	37	54.8
		28820	28838	53.7		4580 4581	25079	25061	52.7	52.6	0.2	662	75.8	41.2	37	54
		27369	27389	52.5			29358	29339	52.8	50	0.9	539	77.1	45.1	37	54.9
1063		7725	7742	50		4582 4583	27468	27451	51.1	50	1.4	100	71.2	. 44	38	50.3
	2084	16549	16567	54.9		4584	8187 17040	8167	50.4	42.9	0.3	463	75.6	41.9	38	53
_		3221	3239	51.5		4585		17021	53.4	50	1.5	492	75.9	42.3	38	54.2
1066		16549	16567	54.9	_	4586	3500	3481	51.2	50	0.3	280	74.6	41.8	38	52.7
1067	2087	20138	20158	50.1		4587	17041 20615	17022	54.1	50	0.8	493	75.8	42.2	38	54.4
		20078	20099	50.5	-	4588	20615	20597 20597	50.6	47.4	0.5	478	75	40.4	38	52.7
1069		13039	13057	51.1		4589	13325	13305	50.6 50.5	47.4	0.1	538	75.3	40.5	38	52.9
		16549	16567	54.9		4590	17041			47.6	0.6	287	75.8	44.6	38	53.3
1071	2091	13701	13725	53.6		4590	14124	17023	53.5	52.6	1.4	493	75.8	42.2	38	54.2
		12975	12993	51.4		4592	13320	14106	52.4 51.4	52.6	1.2	, 424	75.1	41	38	53.4
1073		16548	16566	54.9		4593	16779			47.6	0	346	76.1	44.2	38	53.8
1074		3361	3381	50.5		4594	3500	16758 3481	53.5	50	1.4	232	74	41.4	38	52.9
1075		3361	3381	50.5		4594	3503	3484	51.2 51.5	50	0.7	140	74.1	46.4	38	52.1
1076		16368	16387	50.2		4595 4596	16780	16760	51.5	50 42.9	1	143	74.4	46.9	38	52.3
	2097	7725	7742	50		4597	8188	8168	50.4	42.9	1.2	413	74.9	40.7	38	52.6
1078		8867	8887	52.3	47.6		9597	9573	53.4		0.3	464	75.6	41.8	38	53
1079		2223	2244	51.4		4599	2672			40	1.1	731	75.9	41.2	38	53.9
1080		10242	10265	51.4		4600	10605	2654	50.9	52.6	0.5	450	77	45.3	38	54.3
	2101	8867	8888	52.7		4600	9253	10588 9235	51.1	50	0.2	364	74.5	40.1	38	52.6
1082		3361	3381	50.5		4602	3504		51.6	47.4	1.1	387	75.1	41.3	38	53.2
1083		98	118	50.6		4603		3485	50.4	45	0.1	144	74.3	46.5	38	52.2
1084		12233	12251	51.1	52.6		314	296	50.6	47.4	0	217	75.9	46.5	38	53.4
1085		9926	9944	50.5	52.6		12498	12480	50	47.4	1.1	266	74.8	42.5	38	52.5
1086		3360	3380	51.4	42.9		10455	10434	51.1	40.9	0.6	530	75.3	40.6	38	52.9
1087		9926	9944	50.5			3497	3478	51.3	50	0.1	138	74	46.4	38	52.3
1007	-10/	3320	9944	00.0	52.6	4007	10455	10435	50.5	42.9	0	530	75.3	40.6	38	52.9

1088 2108	10140	10159	52.4	50 4608	10608	10589	51	50	1.4	469	75	40.3	38	52.9
1089 2109	9931	9950	50.2	45 4609	10455	10435	50.5	42.9	0.3	525	75.3	40.6	38	52.8
1090 2110	3219	3238	50.7	50 4610	3500	3481	51.2	50	0.5	282	74.7	41.8	38	52.6
1091 2111	3219	3238	50.7	50 4611	3497	3478	51.3	50	0.6	279	74.7	41.9	38	52.6
1092 2112	3360	3380	51.4	42.9 4612	3500	3481	51.2	50	0.3	141	74	46.1	38	52.3
1093 2113	3360	3380	51.4	42.9 4613	3503	3484	51.5	50	0	144	74.3	46.5	38	52.5
1094 2114	2223	2244	51.4	45.5 4614	2672	2653	51.6	50	0.2	450	77	45.3	38	54.4
1095 2115	9922	9941	51.3	50 4615	10449	10428	51.9	40.9	0.7	528	75.4	40.9	38	53.3
1096 2116	13039	13057	51.1	52.6 4616	13312	13294	51	52.6	0.1	274	75.7	44.5	38	53.4
1097 2117	15951	15973	52.1	43.5 4617	16174	16154	50.4	42.9	1.7	224	73.5	40.6	38	51.7
1098 2118	13176	13196	51.4	47.6 4618	13545	13526	52.9	55	1.5	370	77	46.2	38	54.4
1099 2119	11541	11562	51.5	40.9 4619	11983	11965	53	52.6	1.5	443	75	40.6	38	53.1
1100 2120	2429	2447	50.2	47.4 4620	3056	3038	50.8	52.6	0.6	628	76.3	42.7	38	53.6
1101 2121	11545	11563	50.8	47.4 4621	12258	12238	50.3	42.9	0.5	714	76.2	42.7	38	53.5
1102 2122	8868	8889	50.4	40.9 4622	9245	9226	50.0	45	0.4	378	74.9	41	38	52.6
1103 2123	27361	27380	52.4	55 4623	27466	27448	52.3	52.6	0.1	106	72.5	46.2	38	51.5
1104 2124	8861	8880	50.2	45 4624	9340	9319	50.8	45.5	0.6	480	75.5	41.5	38	53
1105 2125	1784	1802	51.8	52.6 4625	2113	2094	50.0	45.5	1.7	330	76.5	44.2	38	53.3
1106 2126	8868	8889	50.4	40.9 4626	9107	9086	51.6	45.5	1.2	240	74	41.2	38	52
1107 2127	19795	19814	50.4	45 4627	20099	20078	50.5	40.9	0	305	74.4	40.7	38	52.3
1108 2128	26708	26731	54.2	41.7 4628	27347	27324	52.3	41.7	1.9	640	75.6	40.7	38	53.7
1109 2129	19794	19813	50	50 4629	19920	19899	50.2	40.9	0.2	127	72.3	43.3	38	50.7
1110 2130	3031	3051	51.3	52.4 4630	3650	3631	53.1	50	1.8	620	76.5	43.1	38	54
1111 2131	3031	3051	51.3	52.4 4631	3647	3628	50.6	45	0.7	617	76.4	42.9	38	53.8
1112 2132	19794	19813	50	50 4632	19922	19902	50.0	42.9	0.7	129	72.5	43.4	38	50.8
1113 2133	12236	12256	51.2	42.9 4633	12994	12976	50.3	47.4	0.8	759	76.4	42.4	38	53.7
1114 2134	26708	26731	54.2	41.7 4634	27467	27449	52.8	47.4	1.4	760	76.4	41.3	38	54.1
1115 2135	19716	19737	52.2	45.5 4635	19922	19901	51.5	45.5	0.7	207	73.5	41.1	38	52
1116 2136	19715	19735	52.5	47.6 4636	19922	19901	51.5	45.5	0.9	208	73.6	41.3	38	52.1
1117 2137	3360	3379	50.7	45 4637	3503	3484	51.5	50	0.7	144	74.3	46.5	38	52.3
1118 2138	9055	9079	52.8	40 4638	9364	9346	53.9	52.6	1.1	310	75.3	42.9	38	53.7
1119 2139	1782	1801	52.7	50 4639	1881	1861	54.5	52.4	1.8	100	72.4	47	. 38	51.6
1120 2140	26708	26727	50	45 4640	27468	27450	51.9	47.4	1.8	761	75.9	41.3	38	53.3
1121 2141	26708	26727	50	45 4641	27468	27451	51.1	50	1.1	761	75.9	41.3	38	53.3
1122 2142	4593	4613	51.5	47.6 4642	4995	4975	51.7	42.9	0.1	403	76	43.4	38	53.8
1123 2143	19709	19730	51.3	40.9 4643	19930	19911	50.7	50	0.5	222	74	41.9	38	52.1
1124 2144	26421	26441	51.5	42.9 4644	26587	26570	50.2	. 50	1.3	167	72.3	40.1	38	50.8
1125 2145	18979	19000	51.6	45.5 4645	19217	19195	51.7	43.5	0	239	73.5	40.2	38	52.1
1126 2146	18703	18724	53.5	50 4646	19476	19453	53.5	41.7	0	774	75.6	40.3	38	54
1127 2147	4255	4276	51.7	45.5 4647	4708	4690	50.3	47.4	1.4	454	75.1	40.7	38	52.8
1128 2148	3232	3252	51.1	47.6 4648	3503	3484	51.5	50	0.4	272	74.6	41.9	38	52.6
1129 2149	26421	26441	51.5	42.9 4649	26656	26636	51.3	47.6	0.2	236	74.2	41.9	38	52.5
1130 2150	3232	3252	51.1	47.6 4650	3504	3485	50.4	45	0.7	273	74.6	41.8	38	52.4
1131 2151	26421	26441	51.5	42.9 4651	26660	26641	50.2	50	1.3	240	74.2	41.7	38	52.1
1132 2152	26421	26441	51.5	42.9 4652	26683	26665	52.7	52.6	1.2	263	74.8	42.6	38	52.9
1133 2153	26421	26441	51.5	42.9 4653	26686	26669	50.5	50	0.9	266	74.8	42.5	38	52.6
1134 2154	26421	26441	51.5	42.9 4654	26691	26673	51.3	47.4	0.1	271	74.8	42.4	38	52.9
1135 2155	18704	18724	50.8	47.6 4655	19476	19456	50.5	42.9	0.3	773	75.5	40.2	38	53.1
1136 2156	18704	18724	50.8	47.6 4656	19482	19463	50.1	45	0.7	779	75.5	40.2	38	53
1137 2157	942	960	52.1	52.6 4657	1498	1481	51	50	1.1	557	76.9	44.5	38	54.3
1138 2158	942	960	52.1	52.6 4658	1497	1480	50.3	50	1.9	556	77	44.6	38	54.1
	J	- 000	JZ.1	J2.0 7000	1-0/	1400	50.5	- 50	1.0	550	'	77.0	90	J4. I

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	2159	13040	13059	50.9	5	4659	13312	13294	51	52.6	0.1	273	75.6	44.3	38	53.3
1140	2160	18696	18715	51.7	7 5	4660	19476	19453	53.5	41.7	1.8		75.6			
1141	2161	18696	18715	51.7	7 5	4661	19476	19456	50.5	42.9			75.6			
1142	2162	3232	3251	50.3	3 50	4662	3503	3484	_	-		272	74.6		+	
1143	2163	3031	3051	51.3	52.4	4663	3646	3625				616	76.4	42.9	_	
1144	2164	9130	9150	51.3	42.9	4664	9560	9541	50.9		0.4	431	75.3	41.3		
1145	2165	18224	18243	53.1	50	4665	18696	18672			0.4	473				
1146	2166	18224	18243				18696	18673			0.8	473	75.7	42,1		
1147	2167	18225	18243				18697	18679			0.5	473	75.7	42.1	38	54
1148	2168	9130	9150	_			9560	9540	51.6				75.8	42.3		
	2169	8866	8885		45		9252	9235	50.1	42.9 50	0.3	431	75.3	41.3		
1150	2170	9130	9150			1.000	9559	9539			1	387	75.1	41.3		52.7
1151		12267	12290				12501	12480	50.6	42.9	0.7	430	75.3	41.4	38	53
1152		3427	3446		50		3650		53.5	45.5	1	235	74.3	42.1	38	53.2
		3427	3446		50		3648	3631 3628	53.1	50	0.4	224	74.3	42.4	38	52.9
		26039	26058		55	1.0.0	26184	26164	52.3	42.9	0.4	222	74	41:9	38	52.6
	2175	3230	3249		45				52.4	42.9	1.6	146	71.8	40.4	38	51.1
	2176	3230	3249	50.1		4676	3503 3504	3484 3485	51.5	50	1.4	274	74.5	41.6	38	52.3
1157		3427	3446			4677	3504		50.4	45	0.3	275	74.4	41:5	38	52.2
	2178	3429	3449	50.4		4678	3646	3625	52	40.9	0.6	220	74	41:8	38	52.5
1159		3429	3449	50.4		4679		3628	50.6	45	0.2	219	73.9	41.6	38	51.9
1160		8866	8885	51.1	42.9		3646	3625	52	40.9	1.6	218	73.7	41:3	38	51.8
1161		3428	3449	52.8	45.5		9249	9231	50.8	47.4	0.3	384	75.1	41.4	38	52.9
1162		18077	18098				3650	3631	53.1	50	0.3	223	74.2	. 42.2	38	52.8
1163		18078	18098	52.9	50		18696	18672	53.9	40	1	620	76.2	42.4	38	54.3
1164		8866	8885	51.5 51.1	47.6		18696	18673	53.4	41.7	1.9	619	76.2	42.3	38	53.9
1165		3229	3248			4684	9249	9230	51.5	45	0.4	384	75.1	41.4	38	53
1166		12267	12290	50.6		4685	3503	3484	51.5	50	0.8	275	74.6	41.8	38	52.5
1167		8220		54.5	41.7		12495	12476	52.7	45	1.9	229	74.1	41.9	38	52.8
1168		18080	8240	54		4687	8929	8910	54.5	55	0.4	710	75.4	40	38	54.1
1169			18098	51.2		4688	18238	18219	50.3	45	0.9	159	74	44.7	38	52
		18080	18098	51.2		4689	18239	18220	50	45	1.2	160	73.9	44.4	38	51.8
1170		18080	18098	51.2		4690	18697	18679	51.9	52.6	0.7	618	76.3	42.6	38	53.8
1172		18076	18097	53.1		4691	18712	18693	54.8	55	1.7	637	76.3	42.5	38	54.4
1173		8866	8885	51.1		4692	9245	9226	50	45	1.1	380	75	41.1	38	52.6
		943	961	50.3		4693	1498	1481	51	50	0.8	556	76.9	44.4	38	54
1174		18075	18095	50.6		4694	18642	18622	50.5	42.9	0.1	568	76.2	42.6	38	53.6
1175	2195	18075	18095	50.6		4695	18662	18641	50.4	40.9	0.2	588	76.3	42.7	38	53.6
		8866	8885	51.1		4696	9107	9086	51.6	45.5	0.5	242	74.1	41.3	38	52.3
	2197	943	961	50.3		4697	1497	1480	50.3	50	0	555	76.9	44.5	38	54
	2198	7400	7417	50.2		4698	8190	8172	50.3	47.4	0.1	791	76.4	42.2	38	53.6
	2199	13039	13058	51.8		4699	13314	13297	51	50	0.9	276	75.7	44.6	38	53.4
	2200	7725	7743	50.8	47.4		8187	8167	50.4	42.9	0.5	463	75.6	41.9	38	53.1
	2201	18074	18094	51.1		4701	18642	18622	50.5	42.9	0.5	569	76.2	42.5	38	53.6
	2202	25782	25805	52.1		4702	26174	26153	51	40.9	1.1	393	74.8	40.5	38	52.8
	2203	9131	9151	50.4	42.9	4703	9560	9541	50.9	45	0.5	430	75.3	41.4	38	52.9
	2204	25782	25805	52.1		4704	26183	26162	52.8	45.5	0.7	402	74.7	40.3	38	53.1
	2205	9131	9151	50.4	42.9	4705	9560	9540	51.6	42.9	1.2	430	75.3	41.4	38	52.9
1186 2		7725	7743	50.8	47.4	1706	8188	8168	50.4	42.9	0.5	464	75.6	41.8	38	53.1
	2207	985	1004	51.1	50	1707	1494	1476	50.7	47.4	0.4	510	76.5	43.7	38	53.9
	2208	13039	13058	51.8	50	1708	13323	13304	51.11	45	0.7	285	75.8	44.6	38	53.5
1189 2	209	9131	9151	50.4	42.9	709	9559	9539	50.6	42.9	0.3	429	75.3	41.5	38	52.9
									30.0		0.01	120	, 0.0	71.0	30	52.9

1190	2210	12352	12375	52.9	41.7 47	710	12499	12480	51.8	45	1.1	148	73.3	43.9	38	52
1191	2211	3225	3244	52.4	55 47	711	3646	3625	52	40.9	0.4	422	75.4	41.7	38	53.5
1192	2212	25676	25697	51.9	40.9 47		25784	25765	53.3	50	1.4	109	70.3	40.4	38	49.9
1193		25363	25381	51.1		713	25548	25531	51.1	50	0	186	73.7	42.5	38	52
1194		25363	25381	51.1		714	25645	25626	50.8	45	0.4	283	74.2	40.6	38	52.3
1195		18074	18093	50.3	45 47		18642	18622	50.5	42.9	0.2	569	76.2	42.5	38	53.5
1196		3225	3244	52.4		716	3647	3628	50.6	45	1.8	423	75.5	41.8	38	53.1
	2217	3225	3244	52.4	55 47		3650	3631	53.1	50	0.7	426	75.5	42	38	53.7
1198	2218	12352	12375	52.9		718	12494	12476	52.2	47.4	0.6	143	73.2	44.1	38	52
1199		13039	13058	51.8		719	13326	13306	50.7	42.9	1.2	288	75.8	44.4	38	53.4
1200	2220	7617	7636	50.9		720	8188	8169	50.5	45	0.5	572	76.1	42.3	38	53.5
1201	2221	988	1006	52.2		721	1697	1678	50.3	45	2	710	76.9	43.8	38	54
1202	2222	12232	12250	51.9	52.6 47		12739	12719	50.3	42.9	1.7	508	75.8	42.1	38	53.3
1203	2223	12232	12250	51.9	52.6 47		12739	12718	51	40.9	1	508	75.8	42.1	38	53.5
1204	2224	988	1006	52.2	52.6 47		1697	1677	51	42.9	1.2	710	76.9	43.8	38	54.2
1205		8867	8886	50.7		725	9341	9322	51.1	50	0.5	475	75.7	41.9	38	53.3
1206		3223	3242	51.8		726	3650	· 3631	53.1	50	1.3	428	75.6	42.1	38	53.6
1200	2227	8867	8886	50.7		727	9340	9319	50.8	45.5	0.1	474	75.6	41.8	38	53.2
	2228	988	1006	52.2		728	1697	1676	51.7	40.9	0.1	710	76.9	43.8	38	54.4
1200		988	1006	52.2		729	1694	1673	51.7	40.9	0.5	. 707	76.9	43.8	38	54.5
	2230	988	1006	52.2	52.6 4		1494	1476	50.7	47.4	1.5	507	76.5	43.8	. 38	53.9
1211	2231	9931	9950	50.2		731	10455	10434	51.1	40.9	1.5	525	75.3	40.6	38	52.8
1212		3224	3242	50.5	52.6 4		3646	3625	52	40.9	1.5	423	75.4	41.6	38	53
	2233	3224	3242	50.5	52.6 4		3647	3628	50.6	45	0.1	424	75.4	41.7	38	53.1
1214		3016	3036	50.2		734	3187	3166	50.3	45.5	0.1	172	74.6	45.3	38	52.4
1214		24559	24579	52		735	25182	25164	51.4	47.4	0.1	624	74.6	41.8	38	53.7
	2236	1782	1802	53.3	47.6 4		1881	1861	54.5	52.4	1.2	100	72.4	47	38	51.8
1217		7880	7900	50.3		737	8188	8169	50.5	45	0.1	309	74.8	41.7	38	52.6
1218		8861	8880	50.2		738	9248	9229	50.5	45	0.1	388	75	41.7	. 38	52.6
1219		8868	8889	50.4	40.9 4		9312	9293	50.6	45	0.1	445	.75.3	41.3	. 38	52.0
1220		17790	17813	54.3		740	18220	18201	56.1	55	1.8	431	74.9	40.4	. 38	53.8
1221	2241	24569	24590	56.6	54.5 4		25184	25164	55.9	52.4	0.7	616	75.9	41.7	38	55
1222		13176	13196	51.4	47.6 4		13328	13307	51.2	45.5	0.2	153	73.4	43.8	38	51.9
1223		8861	8880	50.2		743	9254	9236	50.6	47.4	0.2	394	74.9	40.9	38	52.6
1224		24622	24643	57.1		744	25400	25377	57.2	50	0.1	779	75.7	40.7	38	55.2
1225		8868	8889	50.4	40.9 4		9256	9237	50.8	45	0.4	389	75.7	41.1	38	52.7
	2246	3361	3381	50.5	42.9 4		3497	3478	51.3	50	0.4	137	74.1	46.7	38	52.1
1227		4593	4613	51.5	47.6 4		4711	4693	50.4	47.4	1.1	119	71.5	40.7	38	50.2
1228		19911	19930	50.7	50 4		20615	20597	50.4	47.4	0.1	705	75.5	40.3	38	53.1
1229		3221	3239	51.5	52.6 4		3497	3478	51.3	50	0.2	277	74.6	41.9	38	52.7
1230		3223	3241	50.2	52.6 4		3504	3485	50.4	45	0.2	282	74.8	42.2	38	52.5
1231		3223	3241	50.2	52.6 4		3503	3484	51.5	50	1.2	281	74.9	42.3	38	52.6
1232		3360	3380	51.4	42.9 4		3504	3485	50.4	45	1:2	145	74.2	46.2	38	52.2
1233		4593	4613	51.5	47.6 4		4711	4692	51.2	45	0.3	119	71.5	40.2	38	50.5
1234		4593	4613	51.5	47.6 4		4710	4691	50.2	45	1.4	118	71.6	42.4	38	50.2
1235		3016	3036	50.2	42.9 4		3186	3165	50.4	40.9	0.2	171	74.4	45	38	52.3
1236		29182	29206	55.4	44.94		29301	29282	55.3	55	0.1	120	73.4	46.7	38	53.1
1237		29183	29206	52.9		757	29306	29287	54.6	55	1.7	124	73.3	46.7	38	52.3
1238		29186	29206	51.3		758	29298	29279	52.6	55	1.3	113	72.8	46	38	51.5
1239		16979	17000	52.6	50 4		17483	17465	54.4	52.6	1.8	505	75.9	42.2	38	54
1240		29182	29205	54.6		760	29298	29279	52.6	55	1.9	117	73.1	46.2	38	52
1240	12200	29102	29200	34.0	41./ 4	100	29296	20219	32.0	25	1.9		10.1	40.2	- 36	52

			17000													
1241		16981	17000	51.3		4761	17111	17090	51.1	40.9	0.2	131	74.5	48.1	38	52.0
	2262	13177	13197	50.3	42.9		13949	13932	51.6		1.3	773	75.8	41	38	53.
	2263	8867	8887	52.3		4763	9252	9234	51.4	52.6	0.9	386	75.1	41.5	38	53.
	2264	24420	24440	50.8		4764	25081	25063	52.4	52.6	1.6	662	75.7	40.9	38	53.
-	2265	7727	7745	50.8		4765	8188	8169	50.5	45	0.4	462	75.6	41.8	38	53.
	2266	2387	2405	51.6		4766	3055	3036	50.6	50	1.1	669	76.7	43.3	38	53.9
	2267	2671	2692	52.1	40.9	4767	3055	3036	50.6	50	1.5	385	74.8	40.5	38	52.0
	2268	29182	29202	51.2		4768	29298	29279	52.6	55	1.4	117	73.1	46.2	38	51.6
	2269	24418	24439	52.9		4769	25081	25063	52.4	52.6	0.5	664	75.7	41.1	38	53.8
	2270	12373	12391	50.8		4770	12992	12974	51.2	52.6	0.4	620	76.5	43.1	_38	53.9
	2271	29179	29199	51.4	42.9	4771	29298	29279	52.6	55	1.2	120	73.4	46.7	38	51.9
	2272	1783	1803	54.2		4772	1882	1861	56	50	1.8	100	72.4	47	38	5
1253		12373	12391	50.8		4773	12498	12480	50	47.4	0.7	126	72.8	44.4	38	- 5
	2274	7728	7746	51.7	52.6	4774	8190	8172	50.3	47.4	1.4	463	75.6	41.9	38	53.
1255	2275	16875	16896	52.2	45.5	4775	17064	17045	51.4	50	0.8	190	74.5	44.2	-38	52.7
	2276	1402	1425	52.8	41.7	4776	2103	2082	52	45.5	0.8	702	76.7	43.3	38	54.4
	2277	28971	28993	51.9	43.5	4777	29358	29339	52.8	50	0.9	388	76.3	. 44.3	38	54.
	2278	24380	24399	55	55	4778	25080	25061	54.1	50	. 1	701	75.9	41.4	38	54.5
1259	2279	24380	24399	55	55	4779	25080	25062	53.5	52.6	1.6	701	75.9	41.4	38	54.3
1260	2280	3168	3189	51	45.5	4780	3497	3478	51.3	50	0.3	330	75.3	42.4	.38	53.
	2281	1402	1426	54.1	40	4781	1626	1602	56.1	44	1.9	225	76.4	47.6	38	54.8
1262	2282	24379	24398	55	55	4782	25080	25061	54.1	50	1	702	75.9	41.3	38	54.4
	2283	24379	24398	55	55	4783	25080	25062	53.5	52.6	1.6	702	75.9	41.3	38	54.3
1264	2284	16875	16895	51.6	47.6	4784	17062	17045	50.2	50	1.4	188	74.4	44.1	38	52.2
	2285	12726	12746	51.3	47.6	4785	12998	12979	50.1	45	1.2	273	75.2	43.2	38	52.7
1266	2286	24378	24397	55	55	4786	24517	24494	53.2	41.7	1.8	140	72.7	42.9	38	51.9
	2287	24378	24397	55	55	4787	25080	25061	54.1	50	1	703	75.9	41.3	38	54.4
	2288	28939	28961	55.2	47.8	4788	29306	29285	56.7	54.5	1.5	368	76.6	45.1	38	55.3
	2289	28940	28961	53.1	45.5	4789	29306	29287	54.6	55	1.5	367	76.5	45	38	54.6
1270	2290	28941	28961	51.6	42.9	4790	29298	29279	52.6	55	1	358	76.3	44.7	38	54
	2291	28178	28200	52	43.5	4791	28284	28265	52.9	50	0.9	107	74.7	51.4	38	53
	2292	28941	28961	51.6	42.9	4792	29358	29339	52.8	50	1.2	418	76.5	44.5	38	54.2
	2293	24378	24397	55	55	4793	25080	25062	53.5	52.6	1.6	703	75.9	41.3	38	54.2
	2294	28938	28960	56.1	47.8	4794	29306	29285	56.7	54.5	0.6	369	76.5	45	38	55.5
1275	2295	12234	12252	50.6	47.4	4795	12498	12480	50	47.4	0.5	265	74.7	42.3	38	52.4
1276	2296	28939	28960	54.7	50	4796	29306	29287	54.6	55	0.1	368	76.6	45.1	38	55,1
	2297	28140	28158	54.1	52.6	4797	28411	28393	52.9	52.6	1.1	272	78.8	52.2	38	56.2
	2298	28941	28960	50.9	45	4798	29298	29279	52.6	55	1.7	358	76.3	44.7	38	53.8
	2299	28140	28158	54.1	52.6	4799	28416	28396	52.4	47.6	1.7	277	78.8	52	38	56
	2300	28941	28960	50.9	45	4800	29358	29339	52.8	50	1.9	418	76.5	44.5	38	53.9
	2301	24179	24200	53.3	40.9	4801	24815	24791	54.5	40	1.2	637	75.8	41.3	38	54.1
1282	2302	28938	28956	50.8	47.4	4802	29298	29279	52.6	55	1.8	361	76.4	44.9	38	53.8
1283	2303	12726	12746	51.3	47.6	4803	12992	12974	51.2	52.6	0.1	267	75.2	43.4	38	53.1
	2304	16874	16893	52.1	50	4804	17062	17045	50.2	50	1.9	189	74.6	44.4	38	52.3
1285	2305	1352	1371	56.1	55	4805	1484	1464	54.3	47.6	1.8	133	74.9	48.9	38	53.8
	2306	11540	11561	53.8	45.5	4806	11983	11965	53	52.6	0.7	444	75.1	40.8	38	53.6
1287	2307	24179	24199	52.7	42.9	4807	24815	24792	53.4	41.7	0.7	637	75.8	41.3	38	53.9
1288	2308	16555	16572	50.3	50	4808	16777	16758	51.5	50	1.2	223	73.6	40.8	38	51.7
1289	2309	24178	24198	52.7	42.9	4809	24815	24791	54.5	40	1.8	638	75.7	41.2	38	53.9
1290	2310	3192	3213	51.8	45.5	4810	3650	3631	53.1	50	1.3	459	75.7	42	38	53.6
1291	2311	3192	3213	51.8	45.5	4811	3647	3628	50.6	45	1.2	456	75.6	41.9	38	53.2
													. 5.5			00.2

1292	2312	24174	24195	52.5	40.9	4812	24815	24792	53.4	41.7	0.9	642	75.8	41.3	38	53.9
1293	2313	16553	16571	53.4	52.6	4813	16780	16760	51.4	42.9	2	228	73.7	40.8	38	52.1
1294	2314	16550	16568	54.1	52.6	4814	17041	17023	53.5	52.6	0.6	492	75.9	42.3	38	54.3
1295	2315	3192	3213	51.8	45.5	4815	3646	3625	52	40.9	0.2	455	75.5	41.8	38	53.5
1296	2316	16551	16568	51.1	50	4816	16777	16758	51.5	50	0.4	227	73.9	41.4	38	52.2
1297	2317	12373	12391	50.8	47.4	4817	12998	12979	50.1	45	0.7	626	76.4	43	38	53.6
1298	2318	28868	28887	50.7	45	4818	29414	29395	50.5	50	0.2	547	77	44.8	38	54.2
1299	2319	24028	24047	53.8	50	4819	24815	24791	54.5	40	0.7	788	76.3	42	38	54.6
1300	2320	2427	2445	52.1	52.6	4820	3056	3038	50.8	52.6	1.3	630	76.4	42.9	38	53.8
1301	2321	28867	28886	53.2	50	4821	29306	29288	53.5	52.6	0.3	440	76.9	45.2	38	54.9
1302	2322	24021	24044	52.8	41.7	4822	24815	24791	54.5	40	1.6	· 795	76.2	41.9	38	54.3
1303	2323	28867	28885	51.5	52.6	4823	29414	29395	50.5	50	0.9	548	77.1	44.9	38	54.2
1304	2324	12369	12388	50.6		4824	13155	13137	52.1	52.6	1.5	787	76.8	43.3	38	54
1305	2325	27368	27392	58.2	48	4825	27467	27443	59.4	48	1.2	100	71.2	. 44	38	52.4
1306	2326	27369	27392	57.2	50	4826	27468	27444	58.4	44	1.2	100	71.2	44	38	52.1
1307	2327	27369	27392	57.2	50	4827	27468	27445	58.1	45.8	0.8	. 100	71.2	- 44	38	52.1
1308	2328	23841	23863	53.7	47.8	4828	24022	24003	55.5	55	1.7	182	74.4	44.5	38	53.3
1309	2329	3192	3213	51.8	45.5	4829	3497	3478	51.3	50	0.5	306	75	. 42.2	38	53
1310	2330	23843	23863	50.3	42.9	4830	24526	24506	50.3	42.9	0	684	76.1	41.8	38	53.4
1311	2331	27366	27389	56.1	45.8	4831	27465	27443	56.4	47.8	0.3	100	71.2	. 44	38	51.8
1312	2332	27366	27389	56.1	45.8	4832	27465	27444	55.6	45.5	0.6	100	71:2	44	38	51.6
1313	2333	27366	27389	56.1	45.8	4833	27465	27445	55.1	47.6	1	100	71.2	. 44	38	51.5
1314	2334	16549	16567	54.9	52.6	4834	16779	16758	53.5	50	1.4	231	74	41.6	38	53
1315	2335	27369	27389	52.5	47.6	4835	27468	27448	53.7	47.6	1.1	100	71.2	·· 44	38	50.7
1316	2336	27369	27389	52.5	47.6	4836	27468	27449	52.6	45	0	100	71.2	. 44	38	50.7
1317	2337	27369	27389	52.5	47.6	4837	27468	27450	51.9	47.4	0.7	100	71.2	44	38	50.5
1318	2338	28654	28672	50.6	52.6	4838	29412	29393	50.3	45	0.2	759	77.9	46.1	38	54.7
1319	2339	2429	2447	50.2	47.4	4839	3053	3034	50.3	50	0.1	625	76.3	42.6	39	53.6
1320	2340	1442	1461	51.6	55	4840	1697	1676	51.7	40.9	0	. 256	75.8	45.3	39	53.7
1321	2341	1442	1461	51.6	55	4841	1697	1677	51	42.9	0.6	256	75.8	45.3	39	53.5
1322	2342	1442	1461	51.6	55	4842	1697	1678	50.3	45	1.3	. 256	75:8	45.3	39	53.3
1323	2343	3214	3233	51.1	50	4843	3504	3485	50.4	45	0.7	291	74.8	41.9	39	52.6
1324	12344	3214	3233	51.1	50	4844	3503	3484	51.5	50	0.4	: 290	74.8	42.1	39	52.8
1325	2345	27374	27392	50.6	47.4	4845	27674	27653	52.5	40.9	1.9	301	74.1	40.2	39	52.2
1326	2346	9930	9949	52.2	50	4846	10670	10649	51.3	40.9	0.9	741	75.8	40.9	39	53.5
1327	7 2347	1442	1461	51.6	55	4847	2103	2083	50.6	42.9	1	662	76.7	43.4	39	53.9
1328	32348	8867	8887	52.3	47.6	4848	9375	9354	50.4	40.9	2	509	75.7	41.8	39	53.2
1329	2349	16367	16386	51.4	50	4849	16775	16755	51.1	42.9	0.3	409	75	40.8	39	52.9
1330	2350	18081	18100	51.7	50	4850	18702	18685	50.2	50	1.5	622	76.2	42.4	39	53.5
1331	1 2351	18083	18102	50.6	45	4851	18702	18685	50.2	50	0.4	620	76.1	42.3	39	53.4
1332	2352	18094	18113	51	50	4852	18702	18685	50.2	50	0.8	609	76.1	42.2	39	53.4
1333	3 2353	8865	8884	50.4	45	4853	9254	9236	50.6	47.4	0.2	390	75	41	39	52.7
1334	4 2354	16367	16386	51.4	50	4854	16774	16754	50.4	42.9	1	408	75	40.9	39	52.7
1335	2355	18008	18028	53	52.4	4855	18220	18202	54.8	52.6	1.9	213	74.4	43.2	39	53.1
1336	2356	27369	27389	52.5	47.6	4856	27674	27653	52.5	40.9	0.1	306	74.3	40.5	39	52.9
1337	7 2357	16367	16386	51.4	50	4857	16774	16753	51.1	40.9	0.3	408	75	40.9	39	52.9
1338	2358	1442	1461	51.6	55	4858	2113	2094	50.1	45	1.5	672	76.7	43.3	39	53.8
1339	2359	7876	7895	51.5	45	4859	8190	8172	50.3	47.4	1.2	315	75.1	42.2	39	52.7
1340	2360	18696	18715	51.7	50	4860	19482	19463	50.1	45	1.7	787	75.6	40.3	39	53
134	1 2361	12370	12388	50.1	47.4	4861	12911	12892	50.5	50	0.4	- 542	76.1	42.4	39	53.4
1342	2 2362	887	905	50.1	47.4	4862	1493	1473	52	47.6	1.9	607	77.1	44.6	39	54.1

	3 2363	16367		51.8	47.6	4863	16774	1 1675	1 53.0	6 41.7	1.8	40	3 75	40.	9 3	E0.0
134		16378	1639	7 50.4	4 4	4864	1711									
134	5 2365	16378	1639	50.4	1 45	4865	1678	1676	1 51.		1			41.		
134	6 2366	1402	142	52.8	3 41.7	4866	150				1.8		1			
134	7 2367	16378	16397	50.4	45	4867	1677		_						-	1
134	8 2368	16378	16397	50.4	45	4868	16775					398				
1349	9 2369	16378	16397	50.4	45	4869	16775				0.6			4	_	
1350	2370	16378	16397	50.4	45		16774				0.0			4		
135	1 2371	16378	16397	50.4	45	4871	16774				0.7	397		41.		
1352	2 2372	16378	16397	50.4	45		16774				1.8		75 75	41.	_	
1353	2373	10250	10274	51.6	40		10608				0.6			41.1	_38	
1354	1 2374	16548	16566	54.9	52.6	1	17112				1.6			40.4		
1355	2375	19709	19730				19922	19902			1.0	214	76.3	42.8		54.5
1356	2376	3218		50.5	45		3504	3485			0.1		73.8	41.6		51.8
1357	2377	3218	3237	50.5			3503	3484		,,,	0.1	287	74.7	41.8		52.5
1358	2378	19709	19730	51.3			19920	19899			1.1	286 212	74.7	42		52.6
1359	2379	1402	1422	50.2	42.9		1501	1480	51.9		1.7	100	73.7	41.5		51.8
1360	2380	1402	1422	50.2	42.9	4880	1501	1481	51.2	42.9	1.1		72	46		50.6
1361	2381	8867	8886	50.7		4881	9249	9230	51.5		0.9	100 383	72	46		50.6
1362	2382	19794	19813	50	50		19928	19908	52	52.4	-		75.2	41.5	39	52.9
1363	2383	8867	8886	50.7		4883	9249	9231	50.8	47.4	0.2	135	72.8	43.7	39	51.1
1364	2384	9927	9945	50.8		4884	10183	10165	51.7	47.4	0.2	383 257	75.2	41.5	39	52.9
1365	2385	27366	27384	52.2	52.6	4885	27566	27546	50.7	47.6	1.5		75.3	44	39	53.1
1366	2386	9927	9945	50.8			10183	10166	50.9	50	0.1	201 257	74.7	44.3	-39	52.6
1367	2387	27366	27384	52.2		4887	27568	27548	50.2	42.9	1.9	203	75.3	44	39	53.1
1368	2388	27366	27384	52.2		4888	27571	27551	51.4	42.9	0.8		74.6	43.8	39	52.4
1369	2389	887	905	50.1		4889	1483	1465	50.5	47.4	0.6	206 597	74.5	43.7	39	52.7
1370	2390	27366	27384	52.2		4890	27579	27558	51.1	40.9	1.1		77	44.6	39	54.1
1371	2391	16549	16567	54.9	52.6		16774	16751	53.6	41.7	1.3	214 226	74.9	44.4	39	52.9
1372	2392	19794	19813	50		4892	19916	19895	50.2	40.9	0.2	123	72.1	41.6	39	53
1373	2393	16551	16568	51.1		4893	17062	17045	50.2	50	0.2	512		43.1	39	50.6
1374	2394	12726	12746	51.3	47.6	4894	13155	13137	52.1	52.6	0.8	430	76.4	42.4	39	53.3
	2395	545	564	50.7		4895	1171	1153	50.4	47.4	0.8	627	78.2	47.2	39	53.9
1376	2396	887	905	50.1	47.4	4896	1483	1464	51.3	45	1.2	597	77		39	54.9
1377		9927	9945	50.8	52.6	4897	10356	10336	52.4	47.6	1.6	430	75.6	44.6 42.1	39	54.1
1378		887	905	50.1	47.4	4898	1481	1463	50.5	47.4	0.4	595	77	44.5	39	53.3
1379	2399	12726	12746	51.3	47.6	4899	12911	12891	51.2	47.6	0.1	186	73.5		39	54.1
1380	2400	19795	19814	50.4	45	4900	19917	19896	50.9	45.5	0.1	123	72.1	41.9	39	51.9
	2401	27361	27380	52.4	55	4901	27566	27546	50.7	47.6	1.7	206	75.1	43.1 45.1	39	50.7
1382	2402	27361	27380	52.4	55	4902	27569	27548	50.9	40.9	1.5	209	74.9		39	52.9
1383	2403	27361	27380	52.4	55	4903	27571	27551	51.4	42.9	1.5	211	75	44.5	39	52.8
1384	2404	8867	8886	50.7	50 4	1904	9256	9237	50.8	45	0.1	390	75.1	44.5	39	53
1385	2405	8373	8391	50.7	47.4	1905	9109	9087	50.5	43.5	0.1	737		41.3	39	52.9
1386	2406	19800	19817	50.4	50/4	1906	19927	19908	52.1	55	1.7	128	75.4	40	39	53
1387	2407	19800	19817	50.4		1907	19924	19905	50.1	50	0.3	128	72.6	43.8	39	51
1388	2408	16553	16571	53.4		1908	16774	16751	53.6	41.7	0.3	222		43.2	39	50.7
1389	2409	2427	2445	52.1		909	3053	3034	50.3	50	1.8	627	73.7 76.4	41	39	52.7
1390 2	2410	887	905	50.1		910	1479	1460	51.6	50	1.5	593		42.7	39	53.6
1391	2411	13177	13197	50.3		911	13321	13301	50.3	42.9	0	145	77.1	44.7	39	54.1
1392 2	2412	8374	8395	52.4	45.5 4		9109	9087	50.5	43.5	1.9	736	73.1 75.4	43.4	39	51.3
1393 2	2413	9926	9944	50.5	52.6 4		10183	10165	51.7	47.4	1.2	258	75.2	40.1	39	53.1
					بالتبت	لت	.5.55	.0100	31.7	77.4	1.2	200	/5.2	43.8	39	52.9

	- 2															
	2414	16562				4914	17056	17035	51.8	45.5	0.1	495	75.8	42	2 39	53.7
1395	1	16562	16581	52.6			17056	17035	51.8	45,5	0.8	495	75.8			
	2416	9926		50.5			10183	10166	50.9	50	0.4	258	75.2	43.8	39	
1397		13177	13197	50.3			13325	13305	50.5	47.6	0.2	: 149	73.3	43.6	39	51.5
1398		10141	10160	51		4918	10356	10336	52.4	47.6	1.4	. 216	73.5	40.7	39	
1399		2823		50.4		4919	3185	3164	51	45.5	0.5	363	75.6	42.7		53.1
1400		19800	19818	52.1	52.6	4920	19916	19895	50.2	40.9	1.9	117	71.7	42.7	39	50.3
1401		8063	8084	51.4	45.5		8189	8170	50.6	50	0.8	127	72.3	43.3		50.9
	2422	985	1008	56.1	50		1485	1465	56	52.4	0	501	76.5	43.7	39	55.4
	2423	985	1008	56.1	50		1485	1466	55.6	55	0.5	501	76.5	43.7	39	55.3
1404		985	1008	56.1	50	4924	1495	1474	55.1	45.5	1	511	76.5	43.6	39	55.2
1405		18017	18036	54.8	55	4925	18231	18209	53.5	47.8	1.3	215	74.5	43.3	39	53.3
	2426	985	1008	56.1	50	4926	1497	1476	56.4	50	0.3	513	76.6	43.9		55.5
1407	2427	13039	13057	51.1	[,] 52.6	4927	13155	13137	52.1	52.6	1	117	73.4	47	39	51.8
1408		985	1008	56.1	50	4928	1498	1478	54.9	47.6	1.2	514	76.5	43.8		55.1
1409	1= :==	988	1006	52.2	52.6	4929	1496	1478	50.4	47.4	1.9	. 509	76.5	43.8		53.8
1410		988	1006	52.2	52.6	4930	1497	1480	50.3	50	2	- 510	76.6	43.9		53.8
1411	2431	19856	19875	50.2	45	4931	20033	20016	50.4	50	0.2	178	74.1	43.8		52
1412		988	1006	52.2	52.6	4932	1498	1481	51	50	1.2	511	76.6	43.8	39	54
	2433	3361	3382	51.9	45.5	4933	3650	3631	53.1	50	1.2	290	75.7	44.1	39	53.6
1414		8867	8888	52.7	45.5	4934	9365	9347	53	52.6	0.3	499	75.8	42.1	. 39	54
	2435	24921	24938	50.4	50	4935	25645	25626	50.8	45	0.4	725	75.5	40.4	39	53.1
	2436	3361	3382	51.9	45.5	4936	3647	3628	50.6	45	1.3	. 287	75.6	43.9	39	53.2
1417	2437	24635	24653	50.5	52.6	4937	25398	25378	51.1	42.9	0.6	764	75.5	40.3	39	53.1
1418	2438	8867	8888	52.7	45.5	4938	9256	9237	50.8	45	1.9	390	75.1	41.3	39	52.9
		18017	18036	54.8	55	4939	18712	18693	54.8	55	0	696	76.4	42.5	39	55
	2440	24633	24651	50.1	52.6	4940	25398	25378	51.1	42.9	0.9	. 766	75.6	40.3	39	53
1421	2441	18011	18032	55.7	54.5	4941	18220	18202	54.8	52.6	0.9	210	74.5	43.3	39	53.7
	2442	18014	18032	51	52.6	4942	18223	18206	51.8	50	0.8	210	74.3	42.9	39	52.4
1423		24630	24648	50.8		4943	25398	25378	51.1	42.9	0.2	769	75.6	40.4	39	53.3
	2444	18014	18032	51	52.6	4944	18231	18210	52.2	45.5	1.2	218	74.5	43.1	39	52.5
	2445	18014	18032	51	52.6	4945	18233	18214	52	50	1.1	220	74.7	43.6	39	52.7
	2446	18014	18032	51		4946	18233	18215	51.3	52.6	0.4	220	74.7	43.6	39	52.7
	2447	18011	18031	54.5	52.4	4947	18220	18201	56.1	55	1.6	210	74.5	43.3	39	. 53.6
	2448	3361	3382	51.9	45.5	4948	3646	3625	52	40.9	0.1	286	75.5	43.7	39	53.5
	2449	4658	4677	50.5	50	4949	5306	5288	52.4	52.6	. 2	649	75.5	40.7	39	53.1
1430		18012	18031	53.2	55	4950	18223	18205	53.3	52.6	0.2	.212	74.5	43.4	39	53.2
	2451	18012	18031	53.2		4951	18712	18693	54.8	55	1.7	701	76.4	42.7	39	54.6
	2452	13040	13059	50.9		4952	13325	13305	50.5	47.6	0.4	.286	75.7	44.4	39	53.3
	2453	8867	8888	52.7		4953	9249	9231	50.8	47.4	1.9	-383	75.2	41.5	39	53
	2454	24179	24198	51		4954	24740	24717	52.5	41.7	1.4	562	76	42.2	39	53.6
	2455	18013	18031	50.6	52.6	4955	18229	18209	50.1	42.9	0.5	217	74.4	42.9	39	52.2
	2456	8865	8884	50.4		4956	9340	9319	50.8	45.5	0.3	.476	75.5	41.6	39	53.1
	2457	24558	24577	50.7	50	4957	24936	24919	51.8	50	1.1	379	75.8	43	39	53.3
	2458	8867	8888	52.7	45.5	4958	9249	9230	51.5	45	1.2	383	75.2	41.5	39	53.2
	2459	26039	26058	54		4959	26753	26733	54	52.4	0.1	715	76	41.5	39	54.5
	2460	26039	26058	54	55	4960	26753	26734	52.6	55	1.4	715	76	41.5	39	54.1
	2461	18009	18028	51.6		4961	18223	18206	51.8	50	0.1	215	74.5	43.3	39	52.7
	2462	24482	24503	51.6	40.9	4962	25080	25062	53.5	52.6	1.8	599	75.5	40.7	39	53.4
	2463	8861	8880	50.2	45	1963	9109	9087	50.5	43.5	0.4	249	73.8	40.6	39	51.8
1444	2464	24483	24503	51	42.9	1964	25086	25069	50.3	50	0.6	.604	75.5	40.7	39	53
																30

144	15 2465	1801	1 1803	0 52.	ol 5	-	1									
	16 2466	2448				5 4965	18220					2 21	74.5	43.	3 39	9 53.1
144		2448					2481					33	5 75.5	4	3 39	53.4
	18 2468	_					2508					60	75.5	40.	8 39	
144		2448				9 4968	25082				0.8	60	1 75.5	40.8	B 39	53
145		2448				9 4969	25085					604	4 75.4	40.0	6 39	
145		2448				9 4970	25086				0	608	75.5	40.7	7 39	
145		1801					18223			50	1.1	213	74.4	43.2	2 39	
		18011					18231				0.7	221	74.7	43.4	39	
145	3 2473	18011					18233				0.9	223	74.9	43.9	39	
145		18011			_		18233			52.6	1.6	223	74.9	43.9	39	
	5 2475 6 2476	24419			_		24815		53.4	41.7	1.2	397	75.9	43.1	39	
		18008					18220	18201	56.1	55	1.6	213	74.4	43.2		
145		24420					24527	24507	51	42.9	0.2	108	70.7	41.7		
	8 2478	12232				4978	12994	12976	50.3	47.4	1.6	763	76.4	42.5		
	9 2479	4644				4979	5306	5288	52.4	52.6	0.1	663	75.6	- 40.9		53.8
146		18009				1.000	18712	18693	54.8	55	1.6	704	76.4	42.6		54.6
146		18010				4981	18223	18205	53.3	52.6	1.5	214	74.4	43		52.7
146		24418	24439				24527	24507	51	42.9	1.9	110	71.3	42.7	39	50.3
146		24418	24439				24815	24792	53.4	41.7	0.5	398	75.9	43.2	39	54.1
146		9351	9370				10017	9999	52.8	52.6	1.6	667	75.7	40.9	39	53.4
	5 2485	18011	18029			4985	18229	18209	50.1	42.9	1.2	219	74.4	42.9	39	52.2
	6 2486	13176	13196			4986	13314	13297	51	50	0.4	139	73	43.9	39	51.5
146		3229	3248	50.6		4987	3497	3478	51.3	50	0.6	269	74.5	41.6	39	52,4
	2488	25772	25793			4988	26182	26161	51.2	40.9	1.2	411	74.7	40.1	39	52.8
	2489	3229	3248	50.6			3500	3481	51.2	50	0.5	272	74.5	41.5	39	52.4
	2490	13176	13196	51.4		4990	13323	13304	51.1	45	0.3	148	73.3	43.9	39	51.8
147		25771	25790	51.1	45	4991	26183	26163	51.7	42.9	0.6	413	74.8	40.4	39	52.8
1472		24418	24436	50	47.4	4992	24526	24506	50.3	42.9	0.3	109	71.4	43.1	39	50.1
	2493	25769	25786	50.3		4993	26182	26161	51.2	40.9	0.9	414	74.8	40.3	39	52.6
	2494	18009	18028	51.6		4994	18231	18210	52.2	45.5	0.6	223	74.7	43.5	39	52.9
	2495	18009	18028	51.6			18233	18214	52	50	0.4	225	74.9	44	39	53
	2496	24418	24436	50	47.4	4996	25082	25064	51.1	52.6	1.1	665	75.8	41.2	39	53.2
1477	1	18009	18028	51.6	55	4997	18233	18215	51.3	52.6	0.3	225	74.9	44	39	53
	2498	24418	24436	50		4998	. 25209	25190	50.6	50	0.6	792	76.2	41.9	39	53.5
	2499	25363	25381	51.1		4999	25650	25631	51.3	45	0.1	288	74.2	40.6	39	52.4
	2500	25363	25381	51.1	52.6		25651	25634	50.4	50	0.7	289	74.3	40.8	39	52.2
1481		25354	25372	50.9	52.6		25548	25531	51.1	50	0.2	195	74.1	43.1	39	52.2
1482		18005	18024	51.1		5002	18223	18206	51.8	50	0.6	219	74.4	42.9	39	52.5
1483		18005	18024	51,1		5003	18231	18210	52.2	45.5	1.1	227	74.6	43.2	39	52.7
	2504	25354	25372	50.9	52.6		25651	25632	52.7	50	1.8	298	74.6	41.3	39	52.6
1485		18005	18024	51.1		5005	18233	18215	51.3	52.6	0.2	229	74.9	43.7	39	52.8
1486		18003	18023	53.5	52.4		18712	18693	54.8	55	1.3	710	76.4	42.7	39	54.7
1487	2507	13176	13196	51.4		5007	13326	13306	50.7	42.9	0.7	151	73.4	43.7	39	51.7
	2508	8868	8889	50.4	40.9		9311	9292	50.7	50	0.3	444	75.4	41.4	39	53
1489	2509	25354	25372	50.9	52.6		25832	25811	52.1	50	1.2	479	75	40.3	39	52.9
1490	2510	8375	8396	51.8		5010	9109	9087	50.5	43.5	1,2	735	75.4	40	39	53
	2511	9918	9938	51.4	47.6		10017	9999	52.8	52.6	1.3	100	72.4	47	39	51.2
1492	2512	8375	8396	51.8	45.5		8933	8916	52.2	50	0.4	559	75.1	40.1	39	53.2
1493	2513	17840	17859	50.8		5013	18632	18611	50.2	40.9	0.6	793	76.1	41.5	39	53.4
	2514	13040	13059	50.9		5014	13155	13138	50.4	50	0.5	116	73.2	46.6	39	51.4
1495	2515	25348	25366	51.2	47.4	5015	25650	25631	51.3	45	0.1	303	74.6	41.3	39	52.7
																J/

	2516	25348	25366	51.2		5016	25651	25634	50.4	50	0.7	304	74.7	41.4	39	52.5
1497	2517	13040	13059	50.9	50	5017	13178	13157	50.4	40.9	0.5	139	73.6	45.3	39	51.7
1498	2518	17792	17813	51.6	40.9	5018	18223	18205	53.3	52.6	1.7	432	74.9	40.3	39	53
1499	2519	25348	25366	51.2	47.4	5019	25832	25811	52.1	50	0.9	485	75.1	40.4	39	53
1500	2520	25348	25366	51.2	47.4	5020	25833	25812	51.4	45.5	0.2	486	75	40.3	39	53
1501	2521	25347	25365	52	52.6	5021	25651	25632	52.7	50	0.7	305	74.8	41.6	39	53
1502	2522	8868	8889	50.4	40.9	5022	9252	9234	51.4	52.6	1	385	75.1	41.3	39	52.8
1503	2523	17793	17813	50	42.9	5023	18229	18209	50.1	42.9	0.1	437	74.9	40.3	39	52.5
1504	2524	17793	17813	50	42.9	5024	18231	18211	50.6	47.6	0.6	439	75	40.5	39	52.6
1505	2525	17793	17813	50	42.9	5025	18234	18216	51	52.6	1	442	75.1	40.7	39	52.7
1506	2526	17793	17813	50		5026	18238	18219	50.3	45	0.2	446	75.1	40.8	39	52.7
1507	2527	17793	17813	50		5027	18239	18220	50	45		447	75.1	40.7	39	52.7
	2528	24180	24199	50.3		5028	24938	24921	50.4	50	0.1	759	75.8	40.8	39	53.2
	2529	25348	25365	50.4		5029	25832	25811	52.1	50	1.7	485	75.1	40.4	39	52.8
	2530	25068	25085	50.3		5030	25182	25164	51.4	47.4	1.1	115	73.3	47	39	51.5
1511		29260	29278	51.3		5031	29414	29395	50.5	50	0.8	155	74.3	45.8	39	52.3
	2532	24179	24198	51		5032	24933	24913	51.1	42.9	0.1	755	75.8	40.9	39	53.5
1513		17790	17811	51.6		5033	18223	18205	53.3	52.6	1.7	434	74.9	40.3	39	53
1514		8063	8084	51.4		5034	8190	8172	50.3	47.4	1.1	128	72.2	43	39	50.8
1515		24178	24197	50.3		5035	24936	24919	51.8	50	1.5	759	75.8	41	39	53.2
	2536	17791	17811	50		5036	18229	18209	50.1	42.9	0.1	439	74.9	40.3	39	52.5
1517	2537	17791	17811	50		5037	18231	18211	50.6	47.6	0.6	441	75	40.6	39	52.6
	2538	24174	24194	50.9		5038	24740	24717	52.5	41.7	1.5	567	76	42.2	39	53.6
1519	2539	24174	24194	50.9		5039	24933	24913	51.1	42.9	0.2	760	75.8	40.9	39	53.4
1520		17791	17811	50		5040	18234	18216	51	52.6	1	444	75.1	40.8	39	52.7
1521		17791	17811	50		5041	18238	18219	50.3	45	0.2	448	75.1	40.8	39	52.7
1522		17791	17811	50		5042	18239	18220	50.5	45	0.2	449	75.1	40.8	39	52.7
1523		24035	24053	52.2		5042	24526	24506	50.3	42.9	1.9	492	75.4	41.3	39	53
1524		24035	24053	52.2		5044	24527	24507	51	42.9	1.2	493	75.4	41.2	39	53.2
	2545	17607	17628	52.3		5045	18231	18209	53.5	47.8	1.2	625	75.2	41.2	39	53.4
	2546	29196	29216	52.5		5046	29358	29339	52.8	50	0.3	163	74.9	46.6	39	53.4
1527	2547	17608	17628	50.9		5047	18231	18211	50.6	47.6	0.4	624	75.2	40.1	39	52.9
1528		8868	8889	50.4		5048	9248	9229	50.0	47.0	0.4	381	75.2	41.2	39	52.9
	2549	17608	17628	50.4		5049	18234	18216	51	52.6	0.3	627	75.3	40.2	39	53.1
	2550	29196	29215	51.8		5050	29358	29339	52.8	52.0	1	163	74.9	46.6	39	53.1
1531		24023	24044	51.4		5051	24527	24508	50.5	45	0.9	505	75.4	40.0	39	53
1532		9409	9428	51.6		5052	9989	9968	50.5	40.9	0.6	581	75.4	40.4	39	53.1
1533		29196	29214	51.1		5053	29358	29339	52.8	50	1.7	163	74.9	46.6	39	52.8
1534		8861	8880	50.2		5054	9257	9238	50.5	45	0.3	397	74.9	40.8	39	52.6
1535		17607	17627	51.6		5055	18231	18209	53.5	47.8	1.9	625	75.2	40.8	39	53.2
	2556	29195	29213	51.9		5056	29358	29339	52.8	50	0.9	164	74.8	46.3	39	53
1537		17608	17627	50.2		5057	18231	18211	50.6	47.6	0.9	624	75.2	40.1	39	52.8
1538		985	1004	51.1		5058	1622	1602	51.6	47.6	0.4	638	77.2	44.7	39	54.4
1539		17608	17627	50.2		5059	18234	18216	51.6	52.6	0.8	627	75.3	44.7	39	52.9
	2560	23841	23860	52.1		5060	24496	24478	50.7	52.6	1.4	656	76.1	40.2	39	53.6
1541		23841	23860	52.1		5061	24498	24479	51.2	52.6	0.9	658	76.1	42.1	39	53.8
1541		3404	3422	50.5		5062	3647	3628	50.6	45	0.9	244	74.6	42.1	39	52.5
1543		9349	9367	51.7		5063	10017	9999	52.8	52.6	1.1	669	75.7	42.6	39	53.6
1543		23841	23859	50.5		5064	24496	24478	52.8	52.6		656	76.1		39	53.6
1544		25068	25085	50.3		5065	25548	25531			0.2			42.1		
				_					51.1	50	0.8	481	75.8	42.2	39	53.3
1546	2566	29186	29205	50.1	40	5066	29414	29395	50.5	50	0.4	229	75.4	45	39	52.9

	7 2567	29182			43.5	5067	29358	29339	52.8	3 50	0.4	177	74.6	45.2	2 39	53.2
	8 2568	23841	23859		52.6	5068	24498	24479	51.2	50	0.7	658		42.1	39	
1549		3404	3422	50.5	47.4	5069	3646	3625	52	40.9	1.5	243		42.4		
	2570	29183		50.4	40.9	5070	29414	29395	50.5	50		232	75.4	44.8		
1551		23841	23859	50.5	52.6	5071	24527	24508	50.5	45	0	687	76.1	41.9		
1552		23838	23857	50.4	50	5072	24093	24075	50.9	52.6		256	75.8	45.3		
1553	2573	23838	23857	50.4	50	5073	24496	24478	50.7			659	76.1	41.9		
1554	2574	23838	23857	50.4	50	5074	24498	24479			0.8	661	76.1	41.9	39	53.5
1555		29181	29201	52.4	47.6	5075	29358	29339	52.8		0.4	178	74.8	45.5	39	53.2
1556	2576	29181	29201	52.4	47.6	5076	29414	29395	50.5		1.9	234	75.6	45.3	- 39	
1557	2577	29180	29200	51.7	42.9	5077	29358	29339	52.8		1.2	179	74.7	45.3	39	53.2 52.9
1558	2578	985	1004	51.1	50	5078	1498	1481	51	50	0.1	514	76.5	43.8		52.9
1559	2579	985	1004	51.1	50		1497	1480	50.3	50	0.8	513	76.6	43.8	39	
1560	2580	8859	8879	50	42.9	5080	9254	9236	50.6		0.6	396	75.0	40.9	39	53.8 52.6
1561	2581	29178	29198	51.4	42.9	5081	29414	29395	50.5		. 0.9	237	75.6	45.1	39	
1562	2582	8859	8879	50	42.9	5082	9340	9319	50.8	45.5	0.7	482	75.5	41.5	39	53.2
1563	2583	16909	16928	50.8	45	5083	17109	17089	50.4	42.9	0.4	201	74.9	· 44.8	39	53 52.7
1564	2584	8794	8813	51.6	45	5084	8919	8901	50.4	47.4	1.2	126	71.8			
1565	2585	8794	8813	51.6	45	5085	8920	8902	52.8	52.6	1.2	127	71.8	42.1	39	50.5
1566	2586	985	1004	51.1		5086	1496	1478	50.4	47.4	0.7	512			39	51
1567	2587	18017	18036	54.8	55	5087	18223	18205	53.3	52.6	1.5	207	76.5	43.8	39	53.8
1568	2588	4593	4613	51.5	47.6	5088	4994	4974	51.2	47.6	0.3	402	74.3	43	39	53.1
1569	2589	18017	18036	54.8	55	5089	18220	18201	56.1	55	1.3	204	76.1	43.5	39	53.7
1570	2590	6155	6174	52.1		5090	6486	6467	50.8	45	1.2	332	74.3	43.1	39	. 53.5
-1571	2591	6158	6178	51.3		5091	6486	6467	50.8	45	0.4	329	74.5	40.7	39	52.5
1572	2592	3232	3251	50.3		5092	3500	3481	51.2	50			74.3	40.1	39	52.4
1573	2593	3232	3251	50.3		5093	3497	3478	51.3	50	0.8	269	74.5	41.6	39	52.3
1574	2594	28523	28544	51.6		5094	29298	29279	52.6		1	266	74.5	41.7	39	52.3
1575	2595	28965	28984	52.9		5095	29358	29339	52.8	55 50	0.1	776	78.4	47.3	39	55.5
1576	2596	8866	8885	51.1		5096	9256	9237	50.8	45		394	76.6	44.9	39	54.6
1577	2597	28518	28538	51.2		5097	28672	28654	50.6	52.6	0.3	391	75.1	41.2	39	52.9
1578	2598	6165	6183	51.2	52.6		6486	6467	50.8	52.6 45	0.7	155	76.7	51.6	39	54
1579	2599	6264	6283	50.4		5099	6483	6463	50.8	42.9	0.3	322 220	74:5	40.7	39	52.5
1580	2600	18074	18093	50.3		5100	18233	18215	51.3	52.6	1		73.8	41.4	39	51.8
1581	2601	6271	6291	51.1	47.6		6483	6463	50.2	42.9		160	73.9	44.4	39	51.9
1582	2602	18074	18093	50.3		5102	18231	18210	52.2	45.5	0.9	213	73.5	40.8	39	51.6
1583	2603	6274	6293	50.1		5103	6483	6463	50.2	45.5	1.9	158	73.5	43.7	39	51.7
1584	2604	18074	18093	50.3		5104	18223	18206	51.8	50	0.1	210	73.5	41	39	51.6
1585	2605	5	23	51.3	52.6		314	296	50.6	47.4	1.5	150	73.2	43.3	39	51.4
1586	2606	6343	6364	50.7	45.5		6486	6467	50.8		0.6	310	76.8	46.5	39	54
	2607	3800	3820	50.6	42.9		4445	4425	50.8	45	0.1	144	71.7	40.3	39	50.5
	2608	7615	7635	51.1	47.6		7821	7798		42.9	0	646	75.4	40.2	39	53
1589	2609	7723	7741	52.2	52.6		8049	8032	52.8 50.4	41.7	1.6	207	73.7	41.5	39	52
1590		1	19	50.1	52.6		314			50	1.8	327	74.9	41.6	39	52.6
	2611	7725	7742	50	50 5		7856	296 7836	50.6	47.4	0.6	314	76.8	46.5	39	53.9
1592		18074	18094	51.1	42.9		18233		51.1	42.9	1.1	132	71.3	40.2	39	50
	2613	3168	3189	51	45.5		3503	18215	51.3	52.6	0.3	160	73.9	44.4	39	52.1
	2614	18074	18094	51.1	42.9 5		18231	3484	51.5	50	0.5	336	75.3	42.6	39	53.1
	2615	13177	13197	50.3		115		18210	52.2	45.5	1.1	158	73.5	43.7	39	51.9
1596		28190	28209	54.2	55 5		13312	13294	51	52.6	0.7	136	72.7	43.4	39	51.1
	2617	28190	28209	54.2	55 5		28671	28652	52.8	55	1.5	482	79.9	52.1	39	56.8
.007	-311	20100	20209	34.2	20 5	11/	28673	28654	53.5	55	0.7	484	79.9	52.3	39	57.1

1598	2619	28190	28208	51.7	52.6 5118	28671	28652	52.8	55	1	482	79.9	52,1	39	56.5
1599		28190	28208	51.7	52.6 5119	28671	28653	50.2	52.6	1.5	482	79.9	52.1	39	56.1
	2620	18074	18094	51.1	42.9 5120	18223	18206	51.8	50	0.7	150	73.2	43.3	39	51.6
	2621	28185	28205	53.5	47.6 5121	28284	28265	52.9	50	0.6	100	74.5	52	39	53.1
	2622	28187	28205	53.1	52.6 5122	28672	28653	51.8	55	1.2	486	79.9	52.3	39	56.6
	2623	2371	2389	50,3	47,4 5123	2900	2881	50.1	45	0.2	530	76.8	44.3	39	53.9
	2624	2371	2389	50.3	47.4 5124	3052	3033	50.3	50	0.1	682	76.7	43.4	39	53.9
	2625	2371	2389	50.3	47.4 5125	3056	3038	50.8	52.6	0.5	686	76.7	43.4	39	53.9
	2626	18074	18095	52.2	45.5 5126	18223	18205	53.3	52.6	1.1	150	73.2	43.3	39	52
1607		2220	2239	51.3	45.5127	2891	2873	50.8	47.4	0.5	672	76.8	43.8	39	54.1
			18097		45 5127	18662	18641	50.4	40.9	1.1	586	76.2	42.7	39	53.6
1608		18077		51.5			28653	50.4	52.6	0.4	555	80	51.9	39	56.1
1609		28117	28135	50.6	52.6 5129	28671 28671	28653	50.2	52.6	0.4	556	79.9	51.8	39	56.1
1610		28116	28134	50.8	47.4 5130		12981	51.1		0.8	769	76.5	42.5	39	56.1
1611		12232	12250	51.9	52.6 5131	13000			45						
1612	2632	18080	18098	51.2	52.6 5132	18702	18685	50.2	50	1	623	76.2	42.4	39	53.5
	2633	12232	12250	51.9	52.6 5133	12999	12980	50.6	40	1.4	768	76.4	42.4	39	53.8
	2634	28820	28840	54.8	47.6 5134	29306	29285	56.7	54.5	1.9	487	77.1 77.1	45.4	39	55.5 55.5
	2635	28820	28840	54.8	47.6 5135	29306	29287	54.6	55	0.3	487		45.4	39	
	2636	18080	18098	51.2	52.6 5136	18642	18622	50.5	42.9	0.7	563	76.2	42.6	39	53.6
	2637	8865	8884	50.4	45 5137	9249	9231	50.8	47.4	0.4	385	75.1	41.3	39	52.8
-	2638	8865	8884	50.4	45 5138	9249	9230 9087	51.5	45 43.5	1.1	385	75.1 73.9	41.3	39 39	52:8
	2639	8865	8884	50.4	45 5139	9109		50.5		0.1	245		40.8		51.9
	2640	28819	28839	56.6	52.4 5140	29306	29285	56.7	54.5	0.2	488	77.2	45.5	39	56.1
1621	2641	9130	9151	52	40.9 5141	9364	9346	53.9	52.6	2	235	74.8	43.4	39	53.1
1622	2642	28820	28839	54.3	50 5142	29306	29287	54.6	55	0.3	487	77.1	45.4	39	55.4
1623		8865	8884	50.4	45 5143	9248	9229	50.1	45	0.3	384	75	41.1	39	52.7
1624		18080	18098	51.2	52.6 5144	18229	18209	50.1	42.9	1.1	150	73.2	43.3	39	51.4
1625		15752	15772	50.8	47.6 5145	16175	16155	51.8	47.6	1	424	75.1	41	39	52.9
	2646	12232	12250	51.9	52.6 5146	12498	12480	50	47.4	1.9	267	74.7	42.3	39	52.4
1627	2647	18078	18098	51.5	47.6 5147	18223	18205	53.3	52.6	1.8	146	73	43.2	39	51.6
	2648	7833	7853	50.7	47.6 5148	8054	8035	50.4	50	0.2	222	74.6	43.2	39	52.4
1629		230	248	51.2	52.6 5149	713	695	50.7	47.4	0.5	484	79.3	50.6	39	55.8
	2650	1472	1491	51.2	45 5150	2153	2134	50.4	45	0.8	682	76.5	42.8	39	53.7
1631		18076	18098	54.4	47.8 5151	18220	18201	56.1	55	1.8	145	73.1	43.4	39	52.6
1632		1442	1461	51.6	55 5152		1673	51.7	40.9	0.1	253	75.9	45.5	39	53.7
1633		28618	28636	52.5	52.6 5153		29339	52.8	50	0.3	741	78.2	47	39	55.6
1634		940	959	56.3	55 5154		1677	54.7	40	1.6	762	77.1	44.2	40	55.5
1635		18076		53.1	45.5 5155		18672	53.9	40	0.8	621	76.2	42.4	40	54.4
1636		940	959	56.3	55 5156		1673	54.4	40	1.9	758	77.2	44.3	40	55.5
1637		3016	3036	50.2	42.9 5157		3167	50.2	40.9	0.1	173	74.5	45.1	40	52.3
1638		18077	18097	51.5	47.6 5158		18673	53.4	41.7	1.9	620	76.2	42.4	40	53.9
1639		18077	18097	51.5	47.6 5159		18679	51.9	52.6	0.3	621	76.2	42.5	40	53.9
1640		9352	9371	50.6	45 5160		9968	51	40.9	0.4		75.4	40.3	40	53
1641		6042	6062	50.4	47.6 5161	6374	6353	50	40.9	0.3		74.6	40.8	40	52.3
1642		942		52.1	52.6 5162		1678	50.3	45	1.8		77.2	44.3	40	54.2
1643		942	960	52.1	52.6 5163		1677	51	42.9	1.1	756	77.2	44.3	40	54.4
1644		6042		50.4	47.6 5164		6273	50.8	45	0.4		73.9	40.6		51.9
1645		942		52.1	52.6 5165		1676	51.7	40.9	0.5		77.2	44.3	40	54.6
1646		942		52.1	52.6 5166		1673	51.7	40.9	0.4		77.2	44.4	40	54.7
1647		13176		51.4	47.6 5167		13727	50.5	43.5	0.9		76.2	42.7	40	53.6
1648	2668	13176	13196	51.4	47.6 5168	13949	13932	51.6	50	0.2	774	75.9	41.1	40	53.6

164	9 2669	942	960	52.1	I FO	6 5169	1 4400	4470	T							
	0 2670	6042				65170	1493	1473		1					40	54.5
165		6042					6292	6272			1.1				40	51.9
165		6042				6 5171	6290	6270	_		0.6			40.6	40	51.9
	3 2673	9402	9420			6 5172	6289	6267	52.2		1.8	248	73.9	40.7	40	51.9
1654		943				4 5173	10017	9999	_		1.4		75.6	41.1	40	53.5
1659		9139		50.3		4 5174	1697	1678			0	755	77.1	44.2	40	54.2
	2676		9159				9324	9300	52.9	40	0.4	186	73.9	43	40	52.6
		9139	9159			5176	9324	9301	52.4	41.7	0.1	186	73.9	43	40	52.6
1657		943	961	50.3		5177	1697	1677	51	42.9	0.7	755	77.1	44.2	40	54.2
1658		6222	6246	52.2		5178	6486	6467	50.8	45	1.4	265	73.8	40	-40	- 52
1659		3895	3914	50.3		5179	4608	4590	51.5	52.6	1.2	714	75.5	40.3	40	53
1660		3889	3911	54.2		5180	4610	4590	53.2	52.4	1.1	722	75.5	40.4	40	53.9
1661		3889	3908	51.3			4608	4590	51.5	52.6	0.3	720	75.5	40.4	40	53.4
1662		9139	9159	52.5	47.6	5182	9359	9335	54.5	40	1.9	221	74.7	43.4	40	53.1
1663		943	961	50.3	47.4	5183	1697	1676	51.7	40.9	1.4	755	77.1	44.2	40	54.2
1664		943	961	50.3	47.4	5184	1694	1673	51.7	40.9	1.5	752	77.2	44.3	40	54.2
1665		6302	6321	51.4	50	5185	6483	6463	50.2	42.9	1.2	182	72.9	40.7	40	51.2
1666		9409	9428	51.6	45	5186	10017	9999	52.8	52.6	1.2	609	75.6	41.1	40	53.5
1667		943	961	50.3	47.4	5187	1493	1473	52	47.6	1.7	551	76.8	44.3	40	54
	2688	13039	13058	51.8	50	5188	13312	13294	51	52.6	0.8	274	75.7	44.5	40	53.4
1669	2689	3799	3820	52.9	45.5	5189	4565	4542	53.9	41.7	1	767	.75.5	40.2	40	53.8
1670		985	1004	51.1	50	5190	1481	1463	50.5	47.4	0.6	497	76.4	43.5	40	53.7
1671		13039	13058	51.8	50	5191	13325	13305	50.5	47.6	1.3	287	75.8	44.6	40	53.7
1672	2692	7615	7635	51.1		5192	8049	8032	50.4	50	0.8	435	75.7	42.3	40	53.2
1673	2693	~ 7615	7635	51.1	47.6	5193	7853	7833	50.7	47.6	0.4	239	74.4	42.3	40	52.4
1674	2694	3034	3053	50.3		5194	3503	3484	51.5	50	1.2	470	76.3	43.4	40	
1675	2695	9140	9159	50.1		5195	9334	9315	52.1	50	2	195	74.3			53.6
1676	2696	3799	3819	51.3		5196	4186	4168	51.8	52.6	0.5	388	75:3	43.6	40 40	52.1
1677	2697	3799	3819	51.3		5197	4434	4416	51.5	52.6	0.3	636	75.3	41.8		53.2
1678	2698	3799	3819	51.3		5198	4435	4417	50.5	52.6	0.8	637			40	53.2
1679	2699	7617	7636	50.9		5199	8190	8172	50.3	47.4	0.6	574	75:4	40.3	40	53
1680	2700	18011	18032	55.7		5200	18443	18424	55.9	55	0.0	433	76.1	42.3	40	53.4
1681	2701	18013	18032	52.2		5201	18696	18672	53.9	40	1.7	684	76.1	43.2	40	55.1
1682	2702	18013	18032	52.2	55	5202	18696	18673	53.4	41.7	1.2	684	76.3	42.4	40	54.2
1683	2703	13177	13197	50.3		5203	13545	13527	50.3	52.6			76.3	42.4	40	54.2
1684	2704	9922	9941	51.3		5204	10455	10434	51.1	40.9	0.1	369	77	46.1	40	54.1
1685	2705	7617	7636	50.9		5205	7853	7833	50.7	47.6		534	75.3	40.6	40	53.1
1686	2706	13177	13197	50.3		5206	13329	13308	50.7	40.9	0.3	237	74.4	42.2	40	52.4
	2707	18014	18032	51	52.6		18238	18219	50.3	40.9	0.2	153	73.2	43.1	40	51.4
1688		18014	18032	51	52.6		18239	18220	50.3	45	0.7	225	74.8	43.6	40	52.5
1689	2709	18014	18032	51		5209	18697	18679	51.9	52.6	0.9	226	74.7	43.4	40	52.4
1690		7708	7730	50.6	43.5		7853	7833	50.7	47.6	0.9	684	76.3	42.4	40	53.8
1691	2711	9140	9159	50.1		5211	9358	9338			0.1	146	71.8	40.4	40	50.6
1692		7723	7741	52.2	52.6		7856	7836	51	42.9	0.9	219	74.4	42.9	40	52.2
1693		988	1006	52.2	52.6		1171		51.1	42.9	1.1	134	71.4	40.3	40	50.4
1694		13177	13197	50.3	42.9		13328	1153	50.4	47.4	1.8	184	73.8	42.9	40	51.9
1695		9935	9955	50.4		5214	10608	13307	51.2	45.5	0.9	152	73.3	43.4	40	51.5
1696		985	1008	56.1		5216		10589	51	50	0.6	674	75.8	41.1	40	53.2
1697		13033	13051	52.1			1484	1463	55.5	50	0.5	500	76.4	43.6	40	55.3
1698		12977	12996	50.2	52.6	5217	13179	13158	50.4	40.9	1.7	147	74.3	46.3	40	52.2
1699		12977	12996	50.2			13320	13300	51.4	47.6	1.1	344	76.1	44.2	40	53.4
1.000	-, 10	12011	12990	30.2	40	5219	13321	13301	50.3	42,9	0.1	345	76	44.1	40	53.4

1700	2720	2823	2844	50.4	45.5	5220	3192	3171	51.9	50	1.5	370	75.7	43	40	53.2
1701	2721	18009	18030	54.6	54.5	5221	18443	18424	55.9		1.4	435	76.1	43.2	40	54.7
1702	2722	12976	12995	51.1	45	5222	13320	13300			0.3	345	76.1	44.3		53.7
1703	2723	1046	1064	51.2	47.4	5223	1531	1512	52.7		1.5	486	76.7	44.2	40	54.1
1704	2724	12976	12995	51.1		5224	13321	13301	50.3		0.8	346	76.1	44.2	40	53.5
1705	2725	12976	12994	50.3	47.4		13320	13300	51.4		1.1	345	76.1	44.3	40	
	2726	12976	12994	50.3	47.4		13321	13301	50.3		1.1	346	76.1		-	53.5
1707		18011	18030	52.9	55		18696	18672	53.9		1	686	76.1	44.2	40	53.5
1708	2728	18011	18030	52.9	55		18696	18673	53.4		0.5	. 686		42.4		54.4
	2729	18011	18030	52.9	55		18697	18679	51.9	52.6	1	687	76.3 76.3	42.4	40	54.4
	2730	9140	9159	50.1	45		9374	9353	50.1	40.9	0	235	74.5	42.5	_40	54.1
	2731	3	23	55.4	52.4		204	185	56.6	55	1.3	202	74.5	42.6	40	52.3
	2732	15255	15273	50.3		5232	15761	15741	51.7	47.6	1.4	507		45	40	54.2
	2733	15255	15273	50.3	52.6		15763	15743	52	47.6	1.7	509	75 75	40	40	52.7
	2734	12965	12985	51.2		5234	13320	13300	51.4	47.6	0.2	356	76.1	40.1	40	52.7
		8373	8391	50.7		5235	9060	9039	50.3	40.9	0.2	688		44.1	40	53.7
	2736	12962	12980	50.7		5236	13320	13300	51.4	47.6	0.4	359	75.4	40.1	40	53
		12938	12957	50.9		5237	13155	13137	52.1	52.6	1.2	218	76.2	44.3	40	53.6
	2738	2671	2692	52.1	40.9		3190	3169	50.7	45.5			75.4	45.4	40	53.2
	2739	2671	2692	52.1	40.9		3192	3171	51.9	50	1.5 0.2	520	75:6	41.5	40	53.2
	2740	12938	12956	50.1		5240	13155	13137	52.1	52.6	0.2	. 522	75.7	41.8	40	53.7
1721	2741	26421	26441	51.5		5241	26592	26574	52.1	52.6	0.9	218	75.4	45.4	40	52.9
		18006	18028	54.5		5242	18443	18424	55.9	55.6		172	72.4	40.1	40	51.2
1723		26421	26441	51.5		5243	26656	26635	52.9	45.5	1.4	438	76:1	43.2	40	54.7
	2744	3055	3074	51.1		5244	3210	3190	50.5		1.4	236	74.2	41.9	40	52.5
		7833	7853	50.7		5245	8189	8170	50.6	47.6 50	0.6	156	74.2	45.5	40	52.2
	2746	26421	26441	51.5		5246	26658	26640	50.8	47.4	0.7	357 238	75.6	42.9	40	53.2
	2747	9131	9151	50.4		5247	9328	9310	51	52.6	0.7	198	74.1	41.6	40	52.2
1728		24921	24938	50.4		5248	25650	25631	51.3	52.6 45	0.7	730	74.3	43.4	40	52.2
1729		24921	24938	50.4		5249	25651	25634	50.4	50	0.9	730	75.5	40.4	40	53.1
1730		9130	9151	52		5250	9324	9301	52.4	41.7	0.5	195	75.6 73.9	40.5	40	53.1
1731		9130	9151	52		5251	9324	9300	52.9	40	1	195	73.9	42.6 42.6	40	52.4 52.4
		8376	8396	50.6		5252	9107	9086	51.6	45.5		732	75.4	42.6	40	
1733		11541	11561	50.9		5253	11727	11708	50.4	45.5	0.5	187	73	40.6	40	53.1
1734	2754	11540	11561	53.8		5254	11984	11966	53	52.6	0.5	445	75.1	40.0	40	51.3 53.6
1735	2755	2371	2389	50.3		5255	2672	2654	50.9	52.6	0.5	302	77.1	47.4	40	
1736		2371	2389	50.3		5256	2998	2977	51.1	40.9	0.8	628	76.7	43.5	40	54.2 53.9
1737		11543	11562	50.4		5257	11727	11708	50.4	45	0.0	185	72.9	40.5	40	51.2
1738	2758	26040	26061	56.4		5258	26589	26567	56.1	47.8	0.4	550	75.1	40.5	40	54.5
1739	2759	11541	11562	51.5		5259	11984	11966	53	52.6	1.5	444	75.1	40.5	40	53.1
1740	2760	7728	7746	51.7		5260	8187	8167	50.4	42.9	1.3	460	75.6	40.5	40	53.2
1741	2761	26040	26061	56.4		5261	26657	26634	54.6	41.7	1.9	618	75.5	40.6	40	54.3
1742	2762	2223	2243	50.2		5262	2675	2656	50.4	50	0.2	453	77	45.3	40	54
1743	2763	2220	2239	51.3		5263	2676	2657	50.7	50	0.5	457	76.9	45.1	40	54.1
1744	2764	11541	11560	50.1		5264	11727	11707	51.1	42.9	1	187	73	40.6	40	51.2
1745	2765	24559	24580	54.2	54.5	5265	25088	25070	54.5	52.6	0.2	530	75.5	41.1	40	54.2
1746	2766	12233	12251	51.1	52.6		12998	12979	50.1	45	1	766	76.5	42.6	40	53.7
1747	2767	12233	12251	51.1	52.6		12412	12392	50	42.9	1.1	180	73.2	41.7	40	51.4
1748	2768	24562	24580	50.1	52.6		25086	25069	50.3	50	0.2	. 525	75.4	41	40	52.9
1749	2769	9931	9950	50.2		5269	10608	10589	51	50	0.8	678	75.8	41.2	40	53.2
1750	2770	24562	24580	50.1	52.6		25209	25188	52	45.5	1.9	648	76.1	42	40	53.4
ــــــــــــــــــــــــــــــــــــــ				1						70.0	7.0	. 070	70.1	42	+0	JJ.4

	1 2771	24562	24580	50.1	52.0	5271	25209	2518	51.4	47.6	1.3	648	76.1	4:	2 40	53.4
175	2 2772	3789	3807	51.8	52.6	5272	4445	442	50.6	42.9	1.2					
175	3 2773	26039	26058	54	5!	5273	26656	2663	53.4					40.6		
175	4 2774	26039	26058	54	5.	5274	26660				1.5			40.5		
175	5 2775	3789	3807	51.8	52.6	5275	4444				1.2			40.5	-	
175	6 2776	24559	24579	52			25086				1.6			41.1		
175	7 2777	12235	12253	50.1	52.6	5277	12999				0.5					
175	8 2778	24559	24579	52		5278	25209	2518			0.5			42.5		
175	9 2779	24559	24579	52			25209	25189			0.6		76.1	42.1	_	
1760	2780	887	905	50.1	47.4		1499	1482			0.6	613	76.1	42.1		
176	1 2781	24558	24577	50.7	50		25182	25164					77.1	44.7		
176	2 2782	26039	26058	54	55		26828	26810		1	0.7 1.2	625	76	41.9	_	
1763	3 2783	8866	8885	51.1	45		9597	9577	50.3	42.9		790	76.4	42.4		
1764	2784	7727	7745	50.8			8049	8032	50.3	42.9 50	0.8	732	75.8	41.1		
1765	2785	13177	13197	50.3	42.9		13747	13726		40.9		323	74.8	41.5		
	2786	887	905	50.1	47.4		1498	1481	51	50	0.4	571	76.2	42.6		
1767		1784	1803	52.5		5287	2103	2083	50.6	42.9	0.9	612 320	77.2	44.8		54.1
1768	2788	12235	12253	50.1	52.6		12498	12480	50.6	47.4	2		76	44.4		53.5
1769		1784	1802	51.8	52.6		2103	2083	50.6	47.4	0.1	264	74.7	42.4		52.4
1770	2790	887	905	50.1		5290	1496	1478	50.6	47.4	1.3	320	76	44.4		53.5
1771	2791	1783	1801	52.9		5291	2153	2133	52.1	47.4	0.3	610	77.1	44.8	_	54.1
1772	2792	887	905	50.1	47.4		1494	1476	50.7	/-	0.9	371	. 76	43.7	40	53.9
1773		3791	3809	51.8		5293	4445	4425	50.7	47.4	0.6	608	77:1	44.7	40	54.1
1774	2794	17813	17832	50.1	45	5294	18506	18488		42.9	1.2	655	75.5	40.5		53.1
1775	2795	3791	3809	51.8		5295	4444	4424	51.2	52.6	1.2	694	75.8	41.2	40	53.2
	2796	17840	17859	50.8	45		18233	18215	50.6	42.9	1.2	654	75.5	40.5	40	53.1
1777		17840	17859	50.8	45	5297	18233		51.3	52.6	0.5	394	74.9	40.9	40	52.8
1778		9930	9949	52.2	50		10356	18214	52	50	1.3	394	74.9	40.9	40	52.8
1779		15211	15230	50.2	45	5299	15949	15930	52.4	47.6	0.2	427	75:6	42.2	40	53.7
1780		98	118	50.6	42.9	5300	253		51.1	45	0.9	739	75.5	40.3	40	53
1781		18004	18023	51.1		5301	18233	233 18214	51.8	47.6	1.2	156	74.5	46.2	40	52.4
1782		24420	24440	50.8	42.9	5302	24936	24919	52	50	0.9	230	75	43.9	40	52.9
1783	2803	24420	24440	50.8		5303	24938		51.8	50	1	517	75.9	42.2	40	53.5
1784		98	118	50.6		5304	254	24921 235	50.4	50	0.4	519	75.8	42	40	53.3
1785		11540	11557	50.4		5305	11727	11707	50	45	0.6	157	74.4	45.9	40	52.2
1786		98	118	50.6		5306	642	622	51.1	42.9	0.7	188	73.1	41	40	51.4
1787	2807	15211	15230	50.2		5307	15595		51.6	47.6	0.9	545	79.1	49.7	40	55.6
1788		15255	15273	50.3	52.6			15576	50.8	45	0.6	385	75.1	41.3	40	52.7
	2809	12373	12391	50.8	47.4		15767 12911	15747	50	42.9	0.3	513	75.1	40.2	40	52.6
1790		18009	18028	51.6		5310	18697	12891	51.2	47.6	0.4	539	76.1	42.5	40	53.6
1791	2811	18009	18028	51.6		5311	18696	18679	51.9	52.6	0.2	689	76.4	42.5	40	54
	2812	18009	18028	51.6		5312	18239	18673	53.4	41.7	1.8	688	76.3	42.4	40	54
	2813	18009	18028	51.6		5313	18239	18220	50	45	1.6	231	74.9	43.7	40	52.5
1794	2814	24417	24436	52.6		5314		18219	50.3	45	1.4	230	75	43.9	40	52.7
	2815	10250	10271	50.6		5314	25079	25061	52.7	52.6	0.1	663	75.8	41.3	40	54
	2816	1356	1375	53.8			10356	10336	52.4	47.6	1.8	107	70.8	42.1	40	49.9
	2817	1356	1375	53.8		5316	1484	1464	54.3	47.6	0.5	129	74.4	48.1	40	53.3
	2818	1356	1375	53.8		5317	1484	1465	53.8	50	0	129	74.4	48.1	40	53.3
	2819	24418	24436			5318	1484	1466	53.1	52.6	0.7	129	74.4	48.1	40	53.1
	2820	18008	18028	50 53	47.4 52.4		24560	24542	50.2	47.4	0.1	143	72.4	42	40	50.8
	2821	25363	25381	51.1		5320	18696	18672	53.9	40	0.9	689	76.3	42.4	40	54.4
1001	-021	2000	20001	31.1	52.6	5321	25646	25627	50.5	45	0.7	284	74.1	40.5	40	52.1

1802	2822	9131	9151	50.4	42 0	5322	9333	9315	52.2	52.6	1.9	203	74.6	43.8	40	52.4
1803		9131	9151	50.4		5323	9374	9353	50.1	40.9	0.3	244	74.6	42.6	40	52.4
1804	2824	24380	24399	55		5324	24582	24560	54.2	52.2	0.9	203	74.4	43.3	40	53.4
1805	2825	19802	19820	53		5325	19921	19900	51.8	45.5	1.2	120	72.4	44.2	40	51.3
1806		3055	3075	51.8		5326	3210	3190	50.5	47.6	1.3	156	74.2	45.5	40	52.2
1807	2827	3055	3075	51.8		5327	3207	3187	50.5	47.6	1.3	153	74.2	45.1	40	52.2
1808		3055	3076	52.4		5328	3210	3190	50.5	47.6	2	156	74.2	45.1	40	52.2
1809		24379	24398	55		5329	24582	24560	54.2	52.2	0.9	204	74.2	43.1	40	53.3
1810		7876	7895	51.5		5330	8054	8035	50.4	50		179				
1811	2831	3055	3076	52.4		5331	3207		50.4		1.1		73.5	42.5	40	51.7
1812	2832	9130	9150	51.3	42.9	5332	9324	3187 9300		47.6	2	153	74	45.1	40	_52
	2832	9130	9150	51.3	42.9	5333	9324		52.9	40	1.6	195	73.9	42.6	40	52.2
								9301	52.4	41.7	1.1	195	73.9	42.6	40	52.2
1814		8794	8813	51.6	45		9324	9301	52.4	41.7	0.8	531	75.7	41.6	40	53.6
1815		8794	8813	51.6		5335	9324	9300	52.9	40	1.3	531	75.7	41.6	40	53.6
_	2836	9130	9150	51.3		5336	9328	9310	51	52.6	0.3	199	74.2	43.2	40	52.4
1817	2837	9130	9150	51.3		5337	9333	9315	52.2	52.6	0.9	204	74.5	43.6	·40	52.6
	2838	24179	24200	53.3		5338	24807	24786	51.7	45.5	1.6	629	75.8	41.3	40	53.7
	2839	4593	4613	51.5		5339	4708	4690	50.3	47.4	1.3	116	71.4	42.2	40	50.2
1820		9130	9150	51.3		5340	9374	9353	50.1	40.9	1.2	245	74.6	42.4	40	52.3
1821	2841	29180	29199	50.1		5341	29412	29393	50.3	45	0.2	233	75.5	45.1	40	53
1822		25348	25366	51.2		5342	25772	25753	51.9	50	0.8	425	74.9	40.5	40	52.9
1823		24179	24200	53.3		5343	24815	24792	53.4	41.7	0.2	637	75.8	41.3	40	54.1
1824		8794	8813	51.6	45		9101	9081	50.5	47.6	1.2	308	74.8	41.6	40	52.6
1825		16861	16880	50.8	50		17056	17035	51.8	45.5	1	196	74.7	44.4	40	52.6
1826		16562	16581	52.6		5346	17038	17021	50.7	50	1.9	477	75.7	41.9	40	53.3
1827		16562	16581	52.6		5347	17039	17022	51.4	50	1.2	478	75.6	41.8	40	53.5
1828	2848	16562	16581	52.6	50	5348	17041	. 17023	53.5	52.6	0.9	480	75.7	41.9	40	53.8
1829		3090	3110	50.3	42.9	5349	3647	3628	50.6	45	0.3	558	76.2	42.7	40	53.5
1830	2850	16562	16580	51.9	52.6	5350	17038	17021	50.7	50	1.2	477	75.7	41.9	40	53.3
1831	2851	16562	16580	51.9	52.6	5351	17039	17022	51.4	50	0.5	478	75.6	41.8	40	53.5
1832	2852	24178	24198	52.7	42.9	5352	24815	24792	53.4	41.7	0.7	638	75.7	41.2	. 40	53.9
1833	2853	16562	16580	51.9	52.6	5353	17041	17023	53.5	52.6	1.6	480	75.7	41.9	40	53.6
1834	2854	24179	24198	51	45	5354	24807	24786	51.7	45.5	0.7	629	75.8	41.3	40	53.5
1835	2855	24179	24198	51	45	5355	24818	24797	51.6	40.9	0.6	640	75.8	41.2	40	53.4
1836		3090	3110	50.3	42.9	5356	3646	3625	52	40.9	1.7	557	76.1	42.5	40	53.5
1837	2857	3089	3110	51.8	45.5	5357	3650	3631	53.1	50	1.3	562	76.3	42.9	40	54
1838	2858	29259	29279	54	52.4	5358	29358	29339	52.8	50	1.1	100	72.4	47	40	51.6
1839	2859	8794	8813	51.6	45	5359	8928	8911	51.9	50	0.2	135	72.2	42.2	40	51.1
1840	2860	24176	24197	52.1		5360	24815	24792	53.4	41.7	1.3	640	75.8	41.2	40	53.8
1841	2861	29259	29277	50.9	52.6	5361	29358	29339	52.8	50	2	100	72.4	47	40	51.1
1842	2862	29257	29276	51.3	50	5362	29358	29339	52.8	50	1.5	102	72.6	47.1	40	51.3
1843	2863	9915	9935	51.8	47.6	5363	10017	9999	52.8	52.6	1	103	72.9	47.6	40	51.6
1844	2864	4639	4659	51.1	47.6	5364	5306	5288	52.4	52.6	1.3	668	75.6	40.9	40	53.4
1845	2865	24178	24197	50.3	40	5365	24807	24786	51.7	45.5	1.4	630	75.8	41.3	40	53.2
1846	2866	28653	28671	50.2	52.6	5366	29414	29395	50.5	50	0.3	762	78	46.2	40	54.7
1847	2867	28653	28671	50.2	52.6	5367	29412	29393	50.3	45	0.1	760	78	46.2	40	54.7
1848	2868	28652	28671	52.8	55	5368	29358	29339	52.8	50	0	707	78	46.4	40	55.5
1849		15752	15772	50.8		5369	16213	16195	50.8	52.6	0	462	75.4	41.3	40	53.1
1850		24178	24197	50.3		5370	24818	24797	51.6	40.9	1.4	641	75.7	41.2	40	53.2
1851		19794	19814	51.7		5371	19909	19885	52.5	40	0.8	116	71.8	43.1	40	50.8
1852		8866	8885	51.1		5372	9341	9322	51.1	50	0	476	75.6	41.8	40	53.4
				<u> </u>		1-0.2			V					71.0		

	185	3 2873	1595	1 1597	3 52.	1 43.	5 5373	1617	1615	5 51.	8 47.0	0.3	3 22	5 73.	7 40.	<u>а</u>	ol 500
	185	4 2874	24174	4 2419	5 52.	5 40.	9 5374	2481									
	185	5 2875	. 8866	888	5 51.	1 4	5 5375	9340	931								
	185	6 2876	15951	1 1597	3 52.	1 43.	5 5376	16169								_	
	185	7 2877	15951	1 1597	4 53.3	41.	5377	16175								_	
	185	8 2878	27437	2745	6 50.2	2 40	5378	27541									
	185	9 2879	15650	1567	4 52.9	40	5379	16210				_					
	186	0 2880	8866	888	5 51.1	4		9334			7				40.		
	186	1 2881	8866	888	5 51.1	45	5381	9310									
	186	2 2882	8866	888	51.1	45		9252								-	00.2
	186	3 2883	3360	3379	50.7			3494							41.		
	186	1 2884	8866	888	51.1	45		9248		1		1			45.	_	
	186	2885	18081	18099	51.2	52.6		18697					617		41.		
	1866	2886	8865	8884				9257	9238		-	0.7	393				
	1867	2887	18081	18099	51.2	52.6		18239	18220			1.2			41		
	1868	2888	18081	18099			5388	18238	18219			0.9			44.7		
	1869	2889	28117	28135				28505	28487	50.2			158		44.9		
	1870	2890	8866	8885		45		9109	9087	50.2		0.4	389	79.5	51.9		
	1871	2891	9055	9079		40		9724	9706			0.6	244	73.9	41		- 52
	1872	2892	3403	3423		47.6		3502	3478			1.5	670	75.4	40.3		53.3
	1873	2893	28855	28874		50		29306	29288	55.8		1.7	100	71.6	45		51.5
i	1874	2894	24173	24194			5394	24815	29288			0.6	452	77.1	45.6		54.9
	1875	2895	3094	3113		50	5395	3647	3628	53.4		0.9	643	75.8	41.2		53.9
	1876		24174	24194		42.9	5396	24807		50.6		0.6	554	76.2	42.8		53.5
ı	1877		28856	28875	52.2		5397	29306	24786	51.7	45.5	0.8	634	75.8	41.3		53.4
1	1878		24174	24194	50.9	42.9	5398		29288	53.5		1.3	451	77.1	45.7	40	54.7
ł	1879		28857	28876	51.7	42.9	5399	24818	24797	51.6		0.7	645	75.8	41.2		53.4
ł	1880		8858	8877	51.2	45	5400	29306	29288	53.5	52.6	1.8	450	77:1	45.6	40	54.6
ł	1881	2901	16553	16571	53.4		5400	9254	9236	50.6		0.6	397	75	41.1	40	52.8
ŀ	_	2902	29197	29219	54.8		5402	16777	16758	51.5	50	1.9	225	73.7	40.9	40	52.1
ł		2903	29198	29219	52.6			29301	29282	55.3	55	0.5	105	73.4	48.6	40	52.9
ŀ	1884	2904	28857	28877	52.3		5403 5404	29306	29288	53.5	52.6	0.9	109	73.3	47.7	40	52.2
ŀ		2905	29199	29219	51.2		5405	29306	29288	53.5	52.6	1.2	450	77.1	45.6	40	54.8
ŀ		2906	3094	3113	50			29298	29280	51.4	52.6	0.2	100	72.4	47	40	51.1
ł	1887	2907	3224	3243	52.3		5406 5407	3646	3625	52	40.9	2	553	76.2	42.7	40	53.4
ŀ	1888		29195	29216	53.8			3650	3631	53.1	50	0.8	427	75.5	41.9	40	53.7
ŀ	1889		28867	28885	51.5		5408	29306	29287	54.6	55	0.8	112	73.6	48.2	40	52.8
ŀ	1890	2910	29196	29216	52.5		5409	29358	29339	52.8	50	1.4	492	76.9	44.9	40	54.4
f	1891	2911	28867	28886	53.2		5410	29298	29279	52.6	55	0.1	103	73.3	48.5	40	52.1
ŀ		2912	3093	3113			5411	29415	29395	53.4	52.4	0.2	549	77.1	45	40	55
H		2913	3225	3243	51.7		5412	3650	3631	53.1	50	1.4	558	76.3	42.8	40	54
ŀ		2914	3225	3243		52.6		3646	3625	52	40.9	1.2	422	75.4	41.7	40	53.1
H		2915	28867		50.9	52.6		3647	3628	50.6	45	0.3	423	75.5	41.8	40	53.1
ŀ	1896		3223	28886	53.2		5415	29306	29287	54.6	55	1.4	440	76.9	45.2	40	54.9
H		2917		3241	50.2	52.6		3500	3481	51.2	50	1	278	74.7	42.1	40	52.5
H		2917	28867	28886	53.2		5417	29298	29279	52.6	55	0.5	432	76.8	45.1	40	54.7
H			24034	24053	53.4		5418	24815	24791	54.5	40	1.1	782	76.3	42.1	40	54.5
۲	1899		3221	3239	51.5	52.6		3650	3631	53.1	50	1.6	430	75.5	41.9	40	53.4
H	1900		18080	18099	53		5420	18696	18673	53.4	41.7	0.5	617	76.2	42.5	40	54.3
H		2921	3095	3116	51.9		5421	3650	3631	53.1	50	1.2	556	76.2	42.8	40	54
H	1902		18080	18099	53	50 5		18696	18672	53.9	40	1	617	76.2	42.5	40	54.3
L	1903	2923	28868	28887	50.7	45 5	423	29298	29279	52.6	55	1.9	431	76.8	45	40	54.1

1904	2924	3218	3238	52.1	47.6	5424	3650	3631	53.1	50	1	433	75.5	41.8	40	53.6
1905	2925	8867	8886	50.7	50	5425	9252	9234	51.4	52.6	0.8	386	75.1	41.5	40	52.9
1906	2926	28867	28887	53.7		5426	29306	29287	54.6	55	0.8	440	76.9	45.2	40	55.1
1907	2927	3218	3237	50.5	45		3497	3478	51.3	50	0.8	280	74.6	41.8	40	52.5
1908	2928	29195	29215	53.2	47.6	5428	29306	29287	54.6	55	1.4	112	73.6	48.2	40	52.6
1909		29196	29215	51.8	50		29298	29279	52.6	55	0.8	. 103	73.3	48.5	40	51.9
1910		3218	3237	50.5	45		3500	3481	51.2	50	0.6	283	74.6	41.7	40	52.5
1911		8867	8886	50.7	50		9245	9226	50	45	0.6	379	74.0	41.2	40	52.6
	2932	28868	28888	51.4	42.9		29298	29279	52.6	55	1.2	431	76.8	41.2	40	54.3
	2933	8867	8886	50.7	50		9107	9086	51.6	45.5	0.9	241	74.1	41.5	40	52.2
	2934	28867	28888	54.3	45.5		29306	29287	54.6	55	0.3	440	76.9	45.2	40	55.2
	2935	19906	19925	50.1	50		20615	20597	50.6	47.4	0.5	710	75.5	40.3	40	53
	2936	16551	16568	51.1	50		16775	16756	50.3	45	0.8	225	73.8	41.3	40	51.9
	2937	8861	8880	50.2	45		9341	9322	51.1	50	0.9	481	75.5	41.6	40	51.9
	2938	16368	16387	50.2	45	5438	. 16781	16761	51.3	47.6	1	414		_		
	2939	3055	3074	51.1	50		3209	3189	50.5	47.6	0.6	155	75 74.1	40.8 45.2	40	52.7 52.1
1920		3217	3236	51.1		5440	3650	3631	53.1	50	2	434	75.5	45.2	40	
1921	2941	28868	28889	52	40.9		29298	29279	52.6	55	0.6	431	76.8	41.9	40	53.3 54.5
1922		28867	28889	54.8	43.5		29306	29287	54.6	55	0.0	440	76.9	45.2	40	
1923		3404	3422	50.5		5443	3503	3484	51.5	50	0.2	100	71.6	45.2	40	55.3
	2944	16368	16387	50.2		5444	16777	16758	51.5	50	1.2	410	71.6	40.7	40	50.4 52.6
1925		24029	24047	52.1		5445	24815	24792	53.4	41.7	1.3	787	76.3	40.7	40	54.1
	2946	16368	16387	50.2		5446	16711	16691	51	42.9	0.8	344	75.1	41.9	40	52.7
1927		28867	28890	55.2	41.7		29306	29287	54.6	55	0.6	440	76.9	45.2	40	
1928		29196	29214	51.1	52.6		29298	29279	52.6	55	1.5	103	73.3	45.2	40	55.3
1929		18488	18507	53.7	55		19224	19200	52.4	40	1.3	737	76	41.5	40	51.7 54
1930		28395	28413	50.2	42.1	5450	28506	28488	50.2	47.4	0	112	74,4	50	40	52.2
1931		16551	16568	51.1		5451	17032	17011	52	45.5	0.9	482	75.8	42.1	40	53.5
1932		28871	28891	50.9		5452	29358	29339	52.8	50	1.9	488	76.9	44.9	40	54.2
	2953	28871	28891	50.9		5453	29298	29280	51.4	52.6	0.5	428	76.8	45.1	40	54.2
1934		28870	28891	52.2		5454	29306	29288	53.5	52.6	1.2	437	76.8	45.1	40	54.6
1935		28868	28891	53.8		5455	29301	29282	55.3	55	1.5	434	76.9	45.2	40	55.1
	2956	3404	3422	50.5		5456	3504	3485	50.4	45	0.1	101	71.5	44.6	40	50.3
1937	2957	29195	29213	51.9		5457	29298	29279	52.6	55	0.7	104	73.1	48.1	40	51.9
1938		28938	28956	50.8		5458	29298	29280	51.4	52.6	0.6	361	76.4	44.9	40	53.8
	2959	18488	18507	53.7	55	5459	19210	19191	52	50	1.7	723	76.4	41.6	40	53.9
	2960	3095	3116	51.9		5460	3647	3628	50.6	45	1.3	553	76.2	42.7	40	53.6
1941	2961	3214	3233	51.1		5461	3497	3478	51.3	50	0.2	284	74.7	41.9	40	52.7
1942		24017	24039	53		5462	24815	24791	54.5	40	1.5	799	76.2	41.9	40	54.4
1943		3095	3116	51.9		5463	3646	3625	52	40.9	0.1	552	76.1	42.6	40	54
1944		18550	18571	50.4		5464	19215	19194	50.2	40.9	0.2	666	75.8	41.1	40	53.2
1945	2965	3214	3233	51.1		5465	3500	3481	51.2	50	0.1	287	74.7	41.8	40	52.7
1946		18586	18603	50.4		5466	19224	19200	52.4	40	1.9	639	75.6	40.8	40	53.1
1947	2967	18586	18603	50.4		5467	19217	19196	50.2	40.9	0.2	632	75.6	41	40	53.1
1948		18586	18603	50.4	44.4	5468	19215	19194	50.2	40.9	0.2	630	75.6	41	40	53.1
1949	2969	18590	18608	50.6	42.1	5469	19224	19200	52.4	40	1.8	635	75.6	40.9	40	53.2
1950	2970	15255	15273	50.3		5470	15767	15746	50.7	40.9	0.4	513	75.1	40.2	40	52.7
1951	2971	28942	28961	50.2		5471	29414	29395	50.5	50	0.3	473	76.8	44.6	40	53.9
1952	2972	18590	18608	50.6		5472	19217	19196	50.2	40.9	0.3	628	75.7	41.1	40	53.1
1953		3055	3074	51.1		5473	3207	3187	50.5	47.6	0.6	153	74	45.1	40	52
	2974	18590	18608	50.6		5474	19215	19194	50.2	40.9	0.3	626	75.7	41.1	40	53.1
		الائتى									<u> </u>	, 020	, 0.7	7111		- 50.1

	2975	18591				5475	19224	19200	52.4	40	0.7	634	75.7	41	40	53.6
1956		18591	18611	51.7		5476	19217	19196	50.2				75.7			
1957		18591	18611	51.7	42.9	5477	19215	19194	50.2		1.4					
	2978	28546	28565	52.2	50	5478	28672	28654	50.6	52.6	1.6		76.5		1	
1959		29191	29210	54.4	55	5479	29415	29395	53.4		1		75.7	45.8		
1960		7880	7900	50.3	42.9	5480	8190	8172	50.3		0		74.9			52.6
1961	2981	3167	3189	51.6	43.5	5481	3650	3631	53.1	50	1.5	1	75.8		40	
1962	2982	28965	28984	52.9	55	5482	29306	29288			0.6	342	76.5	45.3	40	54.5
1963	2983	3166	3188	51.6	43.5	5483	3650	3631	53.1	50	1.5	485	75.8		40	53.7
1964	2984	8867	8887	52.3	47.6	5484	9101	9081	50.5	47.6	1.9	235	74.1	41.7	- 40	52.1
1965	2985	23843	23863	50.3	42.9	5485	24013	23995	50.3		0	171	73.7	43.3	40	51.8
1966	2986	3403	3421	53.1	52.6	5486	3503	3484	51.5		1.7	101	71.9	45.5	40	50.9
1967	2987	16549	16567	54.9	52.6	5487	16777	16756	53.4	45.5	1.5	229	74	41.5	40	52.9
1968		8868	8889	50.4	40.9	5488	9109	9087	50.5	43.5	0.1	242	73.9	40.9	40	51.9
	2989	8861	8880	50.2	45		9311	9292	50.7	50	0.6	451	75.3	41.2	40	52.9
1970	2990	8868	8889	50.4	40.9	5490	9257	9238	50.5	45	0.1	390	75.5	41.2	40	52.9
1971		8868	8889	50.4	40.9	5491	9313	9294	50.4	50	0.1	446	75.4	41.5	40	52.7
	2992	23841	23859	50.5		5492	24013	23995	50.3	47.4	0.1	173	73.4	43.9	40	53
1973	2993	28548	28568	50.5	42.9	5493	28672	28654	50.6	52.6	0.1	125	76.2	52.8	40	53.6
1974	2994	8867	8888	52.7	45.5	5494	9310	9291	51.2	45	1.5	444	75.4	41.4	40	53.2
1975	2995	28968	28988	50.9	47.6	5495	29298	29279	52.6	55	1.8	331	76.2	44.7	40	53.7
1976	2996	19907	19926	52.1	55	5496	20615	20597	50.6	47.4	1.6	709	75.5	40.3	40	53.1
1977	2997	8861	8880	50.2	45	5497	9252	9235	50.1	50	0.1	392	75.5	41.1	40	52.6
1978	2998	19909	19929	50.7	52.4	5498	20615	20597	50.6	47.4	0.2	707	75.5	40.3	40	53.1
1979	2999	3361	3382	51.9		5499	3500	3481	51.2	50	0.7	140	74.1	46.4	40	52.3
1980	3000	18696	18715	51.7	50	5500	18881	18862	50.2	45	1.5	186	74.1	43.5	40	52.3
1981	3001	28968	28989	51.5	45.5	5501	29298	29279	52.6	55	1.1	331	76.2	43.5	40	53.9
1982	3002	19709	19730	51.3	40.9	5502	19923	19903	50.9	47.6	0.4	215	73.9	41.9	40	52.1
1983	3003	3361	3382	51.9	45.5	5503	3497	3478	51.3	50	0.6	137	74.1	46.7	40	52.1
1984	3004	3361	3384	53.7	41.7		3495	3473	51.8	43.5	1.9	135	74	46.7	40	52.5
	3005	19709	19730	51.3	40.9	5505	19924	19905	50.1	50	1.2	216	73.9	41.7	40	51.8
1986	3006	16378	16397	50.4	45	5506	16711	16691	51	42.9	0.6	334	75.2	42.2	40	52.9
1987	3007	3361	3382	51.9	45.5	5507	3504	3485	50.4	45	1.5	144	74.3	46.5	40	52.9
1988	3008	18704	18724	50.8	47.6	5508	19406	19388	50.6	47.4	0.1	703	75.4	40.3	40	53.1
1989	3009	8868	8889	50.4	40.9	5509	9314	9295	51.1	50	0.7	447	75.5	41.6	40	53.1
1990	3010	3361	3382	51.9	45.5	5510	3503	3484	51.5	50	0.5	143	74.4	46.9	40	52.6
1991	3011	19709	19730	51.3		5511	19931	19912	50.9	55	0.4	223	74.4	40.9	40	52.6
	3012	16548	16566	54.9	52.6		16777	16756	53.4	45.5	1.5	230	73.9	41.3	40	52.3
	3013	8868	8889	50.4	40.9	5513	9315	9296	50	45	0.4	448	75.4	41.5	40	52.9
	3014	22321	22341	51.6		5514	22460	22441	50.7	45	0.9	140	71.5	41.5	40	50.3
	3015	29182	29202	51.2	42.9		29412	29393	50.3	45	0.9	231	75.4	45	40	53
1996		22173	22193	51	42.9	5516	22460	22441	50.7	45	0.3	288	74.1	40.3	40	52.1
1997		29181	29201	52.4	47.6		29413	29393	51.1	42.9	1.3	233	75.5	45.1	40	53.3
1998	3018	18704	18724	50.8	47.6		18881	18862	50.2	45	0.5	178	73.8	43.3	40	51.9
	3019	20751	20771	51.3	47.6		21301	21278	51.3	41.7	0.5	551	75.5	43.3	40	53.3
2000	3020	29181	29200	50		5520	29412	29393	50.3	45	0.3	232	75.5	45.3	40	53.3
	3021	20751	20771	51.3	47.6		21304	21283	50.5	40.9	0.8	554	75.5	45.3	40	53.1
2002	3022	29173	29197	54.2		5522	29415	29395	53.4	52.4	0.8	243	75.7	45.3	40	54.1
2003	3023	8867	8888	52.7	45.5		9247	9226	52	45.5	0.7	381	75.7	45.3	40	
	3024	8867	8888	52.7	45.5		9255	9236	51.1	45.5	1.6	389	75	41.1	40	53.2
2005	3025	29178	29198	51.4	42.9		29412	29393	50.3	45	1.1	235	75.5	45.1	40	52.9
									50.0	70		200	70.0	45.1	40	53.1

2006		3163	3185	53.6		5526	3650	3631	53.1	50	0.5	488	75.9	42.4	40	54.2
2007	3027	19800	19817	50.4	50	5527	20033	20016	50.4	50	0	: 234	74.9	43.6	41	52.7
2008	3028	8867	8886	50.7	50	5528	9376	9355	51	40.9	0.3	510	75.7	41.8	41	53.3
2009	3029	19800	19817	50.4	50	5529	19930	19910	50.6	47.6	0.2	131	72.6	43.5	41	51
2010	3030	24418	24439	52.9	45.5	5530	25082	25064	51.1	52.6	1.8	665	75.8	41.2	41	53.5
2011	3031	25771	25790	51.1	45	5531	26182	26161	51.2	40.9	0.1	412	74.8	40.3	41	52.8
2012	3032	12976	12994	50.3	47.4	5532	13326	13306	50.7	42.9	0.3	351	76.1	44.2	41	53.5
2013	3033	12976	12994	50.3	47.4	5533	13328	13307	51.2	45.5	0.9	353	76.1	44.2	41	53.5
2014	3034	2823	2844	50.4	45.5	5534	3500	3481	51.2	50	0.7	678	76.3	42.5	41	53.7
2015	3035	18009	18028	51.6	55	5535	18223	18205	53.3	52.6	1.7	215	74.5	43.3	41	52.7
2016	3036	8223	8240	50.4	50	5536	8933	8916	52.2	50	1.8	711	75.4	40.1	41	53
2017	3037	29180	29199	50.1	40	5537	29414	29395	50.5	50	0.4	235	75.5	45.1	41	53
2018	3038	19800	19817	50.4	50	5538	19925	19906	50.1	50	0.4	126	72.4	43.7	41	50.8
2019	3039	25772	25793	52.4	40.9	5539	26183	26162	52.8	45.5	0.4	412	74.8	40.3	41	53.2
2020	3040	14951	14975	52.2	40	5540	15152	15135	51.4	50	0.8	202	73.4	41.1	41	51.9
2021	3041	2823	2844	50.4	45.5	5541	3503	3484	51.5	50	1	681	76.4	42.6	41	53.7
2022	3042	18075	18095	50.6	47.6	5542	18231	18210	52.2	45.5	1.6	157	73.6	43.9	41	51.8
2023	3043	5	23	51.3	52.6	5543	269	251	51.1	52.6	0.1	265	76.4	46.4	41	53.9
2024	3044	9140	9159	50.1	45	5544	9249	9231	50.8	47.4	0.7	110	71.3	42.7	41	50
2025	3045	24418	24439	52.9	45.5	5545	24815	24791	54.5	40	1.6	398	75.9	43.2	41	54.1
2026	3046	8794	8813	51.6	45	5546	9358	9338	51	42.9	0.6	565	75.8	41.8	41	53.5
2027	3047	24418	24439	52.9	45.5	5547	24807	24786	51.7	45.5	1.2	390	75.9	43.3	41	53.8
2028	3048	2387	2405	51.6	52.6	5548	3186	3165	50.4	40.9	1,2	800	76.9	43.5	41	54.1
2029	3049	24418	24439	52.9	45.5	5549	24527	24506	51.7	40.9	1.2	110	71.3	42.7	41	50.5
2030	3050	24418	24439	52.9	45.5	5550	24517	24494	53.2	41.7	0.3	100	70.8	43	41	50.5
2031	3051	4255	4276	51.7	45.5	5551	4836	4817	51.2	45	0.5	582	75.9	41.9	41	53.6
2032	3052	24420	24440	50.8	42.9	5552	25082	25064	51.1	52.6	0.3	663	75.7	41	41	53.3
2033	3053	8867	8887	52.3	47.6	5553	9250	9232	51.6	47.4	0.8	384	75.1	41.4	41	53.2
2034	3054	14951	14975	52.2	40	5554	15275	15257	50.8	52.6	1.3	325	74.6	40.9	41	52.6
2035	3055	2387	2405	51.6	52.6	5555	3185	3164	51	45.5	0.7	799	76.9	43.6	41	54.2
2036	3056	8865	8884	50.4	45	5556	9252	9235	50.1	50	0.3	388	75.1	41.2	41	52.7
2037	3057	24420	24440	50.8	42.9		24818	24797	51.6	40.9	0.8	399	75.8	42.9	41	53.4
2038	3058	24420	24440	50.8	42.9	5558	24807	24786	51.7	45.5	0.9	388	75.8	43	41	53.4
2039	3059	11541	11560	50.1	45	5559	12110	12090	51.1	42.9	1	570	75.9	41.9	41	53.3
2040	3060	2387	2405	51.6	52.6	5560	2672	2653	51.6	50	0	286	77	47.6	41	54.5
2041	3061	24420	24440	50.8	42.9	5561	24526	24506	50.3	42.9	0.5	107	70.8	42.1	41	49.8
2042	3062	11540	11557	50.4	50	5562	12110	12090	51.1	42.9	0.7	571	76	42	41	53.4
2043	3063	6263	6282	50.9	45	5563	6483	6463	50.2	42.9	0.7	221	73.7	41.2	41	51.8
2044	3064	24418	24440	55	47.8	5564	24815	24791	54.5	40	0.5	398	75.9	43.2	41	54.6
2045	3065	18075	18095	50.6	47.6	5565	18233	18214	52	50	1.4	159	74	44.7	41	52.1
2046	3066	18075	18095	50.6	47.6	5566	18233	18215	51.3	52.6	0.7	159	74	44.7	41	52.1
2047	3067	2429	2447	50.2	47.4	5567	3055	3036	50.6	50	0.4	627	76.3	42.6	41	53.6
2048	3068	19800	19818	52.1	52.6	5568	19917	19896	50.9	45.5	1.2	118	71.9	43.2	41	50.7
2049	3069	24481	24500	50.1	45	5569	24936	24919	51.8	50	1.7	456	75.7	42.1	41	53.1
2050	3070	276	294	50.5	47.4	5570	713	695	50.7	47.4	0.2	438	79.1	50.7	41	55.7
2051	3071	19801	19819	53.2	52.6	5571	19927	19908	52.1	55	1.1	127	72.7	44.1	41	51.6
2052	3072	19801	19819	53.2	52.6		19925	19905	51.4	52.4	1.9	125	72.5	44	41	51.3
2053	3073	3800	3824	53.6	40	5573	4318	4294	54.4	40	0.8	519	75.3	40.8	41	53.9
2054	3074	11540	11557	50.4	50		12258	12238	50.3	42.9	0.2	719	76.2	42	41	53.5
2055	3075	24482	24502	50.3	42.9	5575	24938	24921	50.4	50	0.1	457	75.6	41.8	41	53.1
2056	3076	24482	24502	50.3	42.9	5576	24807	24786	51.7	45.5	1.4	326	75.4	42.9	41	53
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	3077	8867	8888	52.7		5577	9364	9346	53.9	52.6	1.2	498	75.8	42.2	41	54
	3078	24481	24502	51.5		5578	25080	25062	53.5	52.6	2	600	75.5	40.8	41	53.4
	3079	8865	8884	50.4		5579	9107	9086	51.6	45.5	1.2	243	74	41.2	41	52
2060		8867	8888	52.7	45.5		9313	9293	52.1	47.6	0.6	447	75.5	41.6	41	53.5
2061		2427	2445	52.1		5581	3055	3036	50.6	50	1.5	629	76.4	42.8	41	53.7
2062		2823	2844	50.4		5582	3504	3485	50.4	45	0.1	682	76.3	42.5	41	53.7
2063		24483	24503	51	42.9	5583	25085	25068	50.3	50	0.6	603	75.4	40.6	41	53
2064		15255	15273	50.3	52.6		15649	15632	50.1	50	0.2	395	75.1	41.3	41	52.7
2065		24483	24503	51	42.9	5585	25082	25064	51.1	52.6	0.1	600	75.5	40.8	41	53.3
2066	3086	8867	8886	50.7	50	5586	9375	9354	50.4	40.9	0.3	509	75.7	41.8	41	53.2
2067	3087	12976	12994	50.3	47.4	5587	13329	13308	50.5	40.9	0.2	354	76.1	44.1	41	53.4
2068	3088	24483	24503	51	42.9	5588	25081	25063	52.4	52.6	1.4	599	75.5	40.7	41	53.2
	3089	379	398	50.1	45	5589	941	922	50.5	50	0.4	563	78.7	48.8	41	55.2
2070	3090	24483	24503	51	42.9	5590	24936	24919	51.8	50	0.8	454	75.7	42.1	41	53.4
2071	3091	19802	19820	53	52.6	5591	19927	19908	52.1	55	0.8	126	72.8	44.4	41	51.7
2072	3092	9934	9953	50.7	50	5592	10670	10649	51.3	40.9	0.6	737	75.7	40.8	41	53.3
2073	3093	8866	8885	51.1	45	5593	9312	9293	50.6	45	0.5	447	75.4	41.4	41	53
2074	3094	19846	19866	51.2	42.9	5594	20033	20016	50.4	50	0.8	188	74.2	43.6	41	52.2
2075	3095	19848	19867	50.7	45		20033	20016	50.4	50	0.3	186	74.1	43.5	41	52.1
2076	3096	9538	9558	50.9	42.9	5596	10017	9999	52.8	52.6	1.9	480	75.5	41.5	41	53.2
2077	3097	8220	8238	51.5	47.4	5597	8933	8916	52.2	50	0.7	714	75.4	40.1	41	53.3
2078	3098	9140	9159	50.1	45	5598	9249	9232	50	50	0.1	110	71.3	42.7	41	50
2079	3099	12976	12994	50.3	47.4	5599	13332	13312	50.9	47.6	0.6	357	76.2	44.3	41	53.5
2080	3100	15752	15772	50.8	47.6	5600	16174	16154	50.4	42.9	0.4	423	75.1	40.9	41	52.8
2081	3101	18074	18094	51.1	42.9	5601	18232	18212	50.6	47.6	0.5	159	73.7	44	41	51.9
2082	3102	24559	24579	52	52.4	5602	25081	25063	52.4	52.6	0.4	523	75.5	41.1	41	53.5
2083	3103	24559	24579	52	52.4	5603	25079	25061	52.7	52.6	0.7	521	75.5	41.3	41	53.6
2084	3104	3169	3191	52.1	47.8	5604	3650	3631	53.1	50	1	482	75.9	42:3	41	53.8
2085	3105	28117	28135	50.6		5605	28672	28654	50.6	52.6	0.1	556	80	52	41	56.3
2086	3106	1809	1829	50.6		5606	2103	2082	52	45.5	1.5	295	75.4	43.4	41	53
2087	3107	24559	24579	52	52.4	5607	24933	24913	51.1	42.9	0.8	375	75.6	42:7	41	53.4
2088	3108	1809	1829	50.6	42.9	5608	2113	2094	50.1	45	0.4	305	75.4	43.3	41	52.9
2089	3109	28116	28134	50.8		5609	28505	28487	50.2	47.4	0.6	390	79.4	51.8	41	55.8
2090	3110	1808	1828	50.6	42.9	5610	2103	2082	52	45.5	1.5	296	75.5	43.6	41	53.1
2091	3111	15951	15975	53.1	40	5611	16210	16192	54.3	52.6	1.2	260	74.3	41.5	41	53.1
2092	3112	8865	8884	50.4	45	5612	9341	9322	51.1	50	0.7	477	75.6	41.7	41	53.1
2093	3113	15	33	50.7	52.6	5613	642	622	51.6	47.6	0.9	628	79	49.2	41	55.6
2094	3114	8861	8880	50.2		5614	9107	9086	51.6	45.5	1.4	247	73.9	40.9	41	51.9
2095	3115	1808	1828	50.6		5615	2113	2094	50.1	45	0.4	306	75.5	43.5	41	53
2096	3116	24562	24580	50.1		5616	24933	24913	51.1	42.9	1.1	372	75.5	42.5	41	53
2097	3117	28116	28134	50.8	47.4		28672	28654	50.6	52.6	0.2	557	80	51.9	41	56.2
2098	3118	24560	24580	51.3		5618	25081	25063	52.4	52.6	1.1	522	75.4	41	41	53.3
2099	3119	24560	24580	51.3	52.4		25079	25061	52.7	52.6	1.4	520	75.5	41.2	41	53.3
2100	3120	16366	16384	50.3	52.6		16775	16755	51.1	42.9	0.7	410	75.1	41.2	41	52.7
2101	3121	16366	16384	50.3		5621	16774	16754	50.4	42.9	0.1	409	75.1	41.1	41	52.7
2102	3122	24569	24590	56.6		5622	25089	25070	55.8	55	0.1	521	75.4	40.9	41	
2103	3123	24569	24590	56.6		5623	25088	25069	55	50	1.6	520	75.4	40.9	41	54.6 54.3
2104		24567	24590	57.8	54.2		25095	25072	59.3	54.2	1.5	529	75.4	40.8		
2105		24568	24591	58.9		5625	25095	25072	59.3	54.2	0.4	529	75.4	40.8	41	55.2
2106		24568	24591	58.9		5626	25091	25072	59.1	54.5	0.4	528	75.4	40.7	41	55.5
2107		24568	24591	58.9	54.2		25090	25069	58.3	54.5	0.2	523				55.5
التنت	للتتنا			50.5	07.6	-	20000	20009	30.3	34.5	0.0	523	75.4	40.9	41	55.4

2108	3128	16366	16384	50.3	52.6	5628	16774	16753	51.1	40.9	0.8	409	75.1	41.1	41	52.8
2109	3129	1806	1825	51.1	45	5629	2103	2082	52	45.5	-1	298	75.5	43.6	41	53.3
2110	3130	8374	8395	52.4	45.5	5630	9107	9086	51.6	45.5	0.8	734	75.5	40.2	41	53.4
2111	3131	24622	24643	57.1	54.5	5631	24935	24913	56.1	47.8	1	314	74.5	40.8	41	54.1
2112	3132	9130	9150	51.3	42.9	5632	9358	9338	51	42.9	0.3	229	74.5	42.8	41	52.5
2113	3133	12936	12957	53.7	45.5	5633	13530	13511	55.6	55	1.9	595	77.4	45.4	41	55.4
2114	3134	8373	8391	50.7	47.4	5634	9107	9086	51.6	45.5	0.9	735	75.4	40.1	41	53.1
2115	3135	1352	1371	56.1	55	5635	1701	1678	54.3	41.7	1.8	350	76.7	45.7	41	55.1
2116	3136	8867	8886	50.7	50	5636	9342	9323	52.1	50	1.4	476	75.7	42	41	53.3
2117	3137	1352	1371	56.1	55	5637	1701	1677	54.7	40	1.4	350	76.7	45.7	41	55.2
2118	3138	9130	9150	51.3	42.9	5638	9249	9232	50	50	1.3	120	71.7	42.5	41	50.3
2119	3139	16861	16880	50.8	50	5639	17062	17045	50.2	50	0.6	202	74.8	44.6	41	52.5
2120	3140	9130	9150	51.3	42.9	5640	9249	9231	50.8	47.4	0.5	120	.71.7	42.5	41	50.5
2121	3141	9130	9150	51.3	42.9	5641	9249	9230	51.5	45	0.2	120	71.7	42.5	41	50.7
2122	3142	8372	8390	50.7	47.4	5642	9060	9039	50.3	40.9	0.4	689	75.4	40.2	41	53
2123	3143	18074	18093	50.3	45	5643	18232	18212	50.6	47.6	0.3	159	73.7	44	41	51.8
2124	3144	2671	2692	52.1	40.9	5644	3193	3172	52.6	50	0.5	523	75.8	41.9	41	53.8
2125	3145	16562	16581	52.6		5645	17064	17045	51.4	50	1.2	503	75.8	42.1	41	53.6
	3146	2671	2692	52.1		5646	3193	3173	51.4	47.6	0.7	523	75.8	41.9	41	53.6
2127	3147	8372	8390	50.7		5647	9107	9086	51.6	45.5	0.9	736	75.5	40.2	41	53.1
2128		12726	12746	51.3		5648	13321	13301	50.3	42.9	0.9	596	76.7	43.6	41	53.9
2129	3149	8867	8886	50.7		5649	9312	9293	50.6	45	0.1	446	75.4	41.5	41	53
2130		16562	16580	51.9		5650	17062	17045	50.2	50	1.7	501	75.8	42.1	41	53.2
2131	3151	27377	27397	53.4		5651	27674	27653	52.5	40.9	0.9	298	74.3	40.6	41	52.8
2132		16556	16573	50.3		5652	17111	17090	51.1	40.9	0.8	556	76.1	42.4	41	53.5
-	3153	7815	7833	51.5		5653	8531	8512	52	45	0.5	717	75.7	40.7	41	53.5
	3154	3223	3241	50.2		5654	3494	3473	50.4	40.9	0.2	272	74.6	41.9	41	52.4
	3155	8372	8390	50.7		5655	9109	9087	50.5	43.5	0.1	738	75.4	40.1	41	53.1
	3156	3041	3065	57.7		5656	3650	3628	56.3	47.8	1.4	610	76.3	42.8	41	55.4
2137		9569	9591	53		5657	10017	9999	52.8	52.6	0.3	449	75.4	41.4	41	53.7
2138	3158	3041	3065	57.7		5658	3649	3625	56.6	44	1.2	609	76.3	42.7	41	55.5
2139	3159	13176	13196	51.4		5659	13321	13301	50.3	42.9	1	146	73.3	43.8	41	51.5
2140	3160	16366	16385	52.9	55	5660	16775	16755	51.1	42.9	1.8	410	75.1	41	41	53
2141	3161	16366	16385	52.9	55	5661	16775	16754	51.7	40.9	1.1	410	75.1	41	41	53.2
2142	3162	16366	16385	52.9	55	5662	16774	16753	51.1	40.9	1.8	409	75.1	41.1	41	53
2143	3163	1402	1422	50.2	42.9	5663	2104	2084	50.6	42.9	0.4	703	76.7	43.2	41	53.8
2144	3164	1402	1422	50.2	42.9	5664	1697	1678	50.3	45	0.1	296	76	44.9	41	53.4
2145	3165	3055	3076	52.4	45.5	5665	3503	3484	51.5	50	1	449	76.1	43.2	41	53.8
2146	3166	15211	15230	50.2	45	5666	16001	15980	51.1	45.5	0.9	791	75.6	40.3	41	53.1
2147	3167	12267	12290	54.5	41.7	5667	12414	12392	53.9	43.5	0.6	148	72.2	41.2	41	51.8
2148	3168	8861	8880	50.2		5668	9256	9237	50.8	45	0.6	396	75	40.9	41	52.6
2149	3169	3049	3071	56.3	52.2	5669	3650	3628	56.3	47.8	0	602	76.4	42.9	41	55.4
2150	3170	3049	3071	56.3	52.2	5670	3648	3625	55.5	41.7	0.8	600	76.3	42.7	41	55.2
2151	3171	8861	8880	50.2	45	5671	9313	9294	50.4	50	0.3	453	75.3	41.3	41	52.9
2152	3172	12352	12375	52.9	41.7	5672	12911	12891	51.2	47.6	1.7	560	76.1	42.5	41	53.7
2153	3173	7965	7985	51.9	42.9	5673	8531	8512	52	45	0.2	567	75.1	40	41	53.2
2154	3174	18017	18036	54.8	55	5674	18233	18212	53.5	50	1.3	217	74.7	43.8	41	53.5
2155	3175	8867	8886	50.7		5675	9257	9238	50.5	45	0.2	391	75.1	41.2	41	52.8
	3176	3221	3239	51.5		5676	3494	3473	50.4	40.9	1.1	274	74.5	41.6	41	52.4
2157	3177	1402	1422	50.2	42.9	5677	1697	1677	51	42.9	0.8	296	76	44.9	41	53.4
	3178	1402	1422	50.2		5678	1697	1676	51.7	40.9	1.5	296	76	44.9	41	53.4
	3179	18011	18032	55.7		5679	18220	18201	56.1	55	0.4	210	74.5	43.3	41	53.9
	1		10002	55.7	07.0	20,0	200	10201	30.1		0.4	210	74.5	40.0	41	55.9

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216	0 3180	1272	6 1274	16 51	.3 47	6 5680	1332	9 1330	08 50	.5 40	9 0.	.8 60	11 70	<u> </u>			_
	1 3181		2 142	52.			169						-				3.9
	2 3182		3 1803	2 52.	.2 5	5 5682	1822							6 44	_		3.8
216	3 3183	377	7 379	7 51.	7 47	6 5683	444							_		41 52	_
216	4 3184	377	7 379	7 51.		6 5684	444							-			3.2
	5 3185	787	6 789	5 51.		5 5685	818			_	0 0.					41 53	_
	6 3186	1801	4 1803	2 5		6 5686	1822				-					41 52	_
216	7 3187	140	2 142	5 52.			169		-	1 42		_				41 52	
216	8 3188	140	2 142	5 52.			169				-		-			41 53	
216	9 3189	140	2 142	5 52.	8 41.		150							1		11 53	
2170	3190	1236	1238	4 51.			1315									11 50	
217	1 3191	2736	2738	0 52.4			27573								_		54
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2174	3194	27361					27567							-		1 53	
2175	3195	9131	915	50.4			9249					_					3
2176	3196	3217	3236	51.1	50	5696	3504							_			
2177	3197	18011	18029	51.3		5697	18232										
2178	3198	3055	3074		_	5698	3503	10-11									
2179	3199	8866	8886	52.3		5699	9364	9346						43.	_		
2180	3200	16368	16387	50.2			16774	16752			_	_	1	42.			
2181	3201	8859	8879			5701	9252	9235			_	-		40.8	-		7
2182	3202	9131	9151	50.4		5702	9249	9231	50.8			394		41.		1	6
2183	3203	3217	3236			5703	3494	3473				119		42.9			
2184	3204	8859	8879	50		5704	9341	9322		1		278	1	41.7			4
· 2185	3205	9131	9151	50.4		5705	9249	9232	51.1			483		41.6		1 5	3
2186	3206	8867	8886	50.7		5706	9248	9232	50			119		42.9	_	1 00.	
2187	3207	27366	27384	52.2		5707	27576	27555	50.1	45	0.5	382	75.1	41.4	4		7
2188	3208	1442	1461	51.6		5708	1879	1861	51	40.9	1.2	211	74.8	44.1	4		
2189	3209	12366	12384	51.7		5709	12724	12705	53 52.4		1.4	438	76.2	43.6	41		
2190	3210	12366	12384	51.7		5710	12498	12480	52.4	55 47.4	0.7	359	75.6	42.9	41		
2191	3211	98	118	50.6	42.9		713	695	50.7	47.4	1.6	133	73	44.4	41	51.2	
2192	3212	12373	12391	50.8	47.4		13155	13138	50.7		0.1	616	79	49.4	41	55.6	
2193	3213	18011	18030	52.9		5713	18230	18209	51.3	50 45.5	0.4	783	76.8	43.4	41	54	1
2194	3214	1402	1426	54.1		5714	1700	1676	53.9	45.5	1.6 0.2	220	74.5	43.2	41	52.7	
	3215	18011	18030	52.9		5715	18223	18205	53.3	52.6		299	76	44.8	41	54.5	ı
2196		1402	1426	54.1		5716	1698	1677	52.3	40.9	0.5	213	74.4	43.2	41	53.1	ı
2197	3217	16463	16483	51.3		5717	17032	17011	52.3	45.5	1.8 0.7	297	76	44.8	41	54	
2198		18009	18030	54.6	54.5		18220	18201	56.1	45.5		570	76	42.1	41	53:7	
2199	3219	1402	1426	54.1		5719	1700	1678	52.9	43.5	1.6	212	74.5	43.4	41	53.6	
	3220	9131	9151	50.4	42.9		9358	9338	52.9	43.5	1.3	299	76	44.8	41	54.2	
	3221	3055	3075	51.8	47.6		3503	3484	51.5	42.9 50	0.6	228	74.6	43	41	52.4	
2202	3222	18013	18031	50.6	52.6		18232	18212	50.6		0.3	449	76.1	43.2	41	53.8	
2203	3223	8794	8813	51.6		723	9249	9230	51.5	47.6 45	9	220	74.7	43.6	41	52.6	
	3224	8794	8813	51.6	45		9249	9231	50.8	47.4	0.1	456	75.4	41.4	41	53.4	
2205 3	225	16549	16567	54.9	52.6		17065	17045	53.1		0.8	456	75.4	41.4	41	53.1	
2206 3	226	8794	8813	51.6	45 5		9249	9232	50	47.6	1.9	517	76	42.4	41	54.2	
2207 3	227	1402	1426	54.1	40 5		2104	2082	53.5	50 43.5	1.6	456	75.4	41.4	41	52.9	
	228	9927	9946	51.3		728	10356	10336	52.4	43.5	0.6	703	76.7	43.2	41	54.8	
2209 3		3219	3238	50.7	50 5		3494	3473	50.4		1.1	430	75.6	42.1	41	53.4	
2210 3	230	16549	16567	54.9	52.6 5		17033	17011	53.2	40.9	0.3	276	74.5	41.7	41	52.4	
						. 50	.,,,,,,	17011	33.2	43.5	1.7	485	75.8	42.1	41	54.1	

	3231	18014	18032	51		5731	18702	18685	50.2	50	0.8	. 689	76.2	42.2	41	53.5
2212		8794	8813	51.6	45	5732	9333	9315	52.2	52.6	0.6	• 540	75.9	42	41	53.7
2213		8867	8888	52.7		5733	9249	9229	53	47.6	0.2	383	75.2	41.5	41	53.5
2214	3234	18009	18028	51.6	55	5734	18229	18209	50.1	42.9	1.5	. 221	74.5	43	41	52.3
2215		9633	9651	51	47.4	5735	10017	9999	52.8	52.6	1.8	385	75.6	42.6	41	53.3
2216	3236	9915	9935	51.8	47.6	5736	10449	10428	51.9	40.9	0.1	535	75.4	40.9	41	53.4
2217	3237	29259	29277	50.9	52.6	5737	29414	29395	50.5	50	0.3	156	74.5	46.2	41	52.4
2218	3238	8868	8889	50.4	40.9	5738	9317	9297	50.5	42.9	0.1	450	75.4	41.6	41	53
2219	3239	29257	29276	51.3	50	5739	29414	29395	50.5	50	0.8	158	74.6	46.2	41	52.5
2220	3240	13176	13196	51.4	47.6	5740	13332	13312	50.9	47.6	0.5	157	73.6	43.9	41	51.9
2221	3241	9918	9938	51.4			10449	10428	51.9	40.9	0.5	532	75.4	40.8	41	53.3
2222	3242	13176	13196	51.4	47.6	5742	13856	13835	50.1	45.5	1.3	681	75.8	41.1	41	53.2
2223	3243	29253	29270	50			29414	29395	50.5	50	0.5	162	75.2	47.5	41	52.8
2224	3244	13037	13058	54.8			13530	13511	55.6	55	0.8	494	77.3	45.7	41	55.6
2225	3245	18009	18028	51.6	55	5745	18702	18685	50.2	50	1.5	694	76.3	42.4	41	53.6
2226	3246	24178	24197	50.3		5746	24938	24921	50.4	50	0.1	761	75.8	40.9	41	53.2
2227	3247	24174	24195	52.5		5747	24740	24717	52.5	41.7	0	567	76	42.2	41	54
2228	3248	7679	7698	50.6	50	5748	8054	8035	50.4	50	0.1	376	75.6	42.6	41	53.1
2229	3249	18005	18024	51.1	50	5749	18229	18209	50.1	42.9	1	225	74.4	42.7	41	52.2
2230	3250	24174	24195	52.5		5750	24933	24913	51.1	42.9	1.4	760	75.8	40.9	41	53.5
2231	3251	3016	3036	50.2	42.9		3500	3481	51.2	50	0.9	485	76.3	43.3	41	53.6
2232	3252	28820	28838	53.7		5752	29306	29288	53.5	52.6	0.2	487	77.1	45.4	41	55.1
2233	3253	18005	18024	51.1		5753	18233	18214	52	50	0.9	229	74.9	43.7	41	52.8
2234	3254	3016	3036	50.2		5754	3503	3484	51.5	50	1.2	488	76.3	43.4	41	53.6
2235	3255	7723	7741	52.2	52.6		8054	8035	50.4	50	1.7	332	75	41.9	41	52.8
2236	3256	29200	29224	54.2		5756	29358	29339	52.8	50	1.4	159	74.5	45.9	41	53.1
2237	3257	3016	3036	50.2	42.9		3504	3485	50.4	45	0.1	489	76.3	43.4	41	53.6
2238	3258	985	1004	51.1	50	5758	1499	1482	50.1	50	1.1	515	76.5	43.7	41	53.7
2239	3259	8866	8885	51.1	45	5759	9257	9238	50.5	45	0.6	392	75	41.1	41	52.8
2240	3260	3016	3036	50.2	42.9	5760	3647	3628	50.6	45	0.4	632	76.4	: 42.9	41	53.7
2241	3261	13039	13058	51.8	50	5761	13749	13727	50.5	43.5	1.3	711	76.7	43.2	41	53.9
2242	3262	24096	24119	54.4	41.7	5762	24815	24792	53.4	41.7	1	720	75.8	• 41	41	54.2
2243	3263	17840	17859	50.8	45	5763	18229	18209	50.1	42.9	0.7	390	74.7	40.3	41	52.4
2244	3264	15255	15273	50.3	52.6	5764	15647	15628	51	45	0.7	393	75.1	41.2	41	52.7
2245	3265	988	1006	52.2	52.6	5765	1500	1482	50.6	47.4	1.6	513	76.5	43.7	41	53.8
2246	3266	24035	24053	52.2	52.6	5766	24527	24508	50.5	45	1.7	493	75.4	41.2	41	53
2247	3267	18616	18636	51.4	47.6	5767	19215	19194	50.2	40.9	1.1	600	75.7	41.2	41	53.1
2248	3268	8374	8393	51.2	45	5768	9101	9081	50.5	47.6	0.7	728	75.5	40.2	41	53.1
2249	3269	17840	17859	50.8	45	5769	18238	18219	. 50.3	45	0.5	399	75	40.9	41	52.7
2250	3270	24030	24047	50.7	50	5770	24526	24506	50.3	42.9	0.4	497	75.5	41.2	41	53
2251	3271	24030	24047	50.7	50	5771	24527	24507	51	42.9	0.3	498	75.4	41.2	41	53.1
2252	3272	17840	17859	50.8	45	5772	18239	18220	50	45	0.8	400	74.9	40.8	41	52.6
2253	3273	985	1008	56.1		5773	1626	1602	56.1	44	0	642	77.1	44.5	41	55.9
2254	3274	13039	13057	51.1	52.6	5774	13749	13727	50.5	43.5	0.6	711	76.7	43.2	41	53.9
2255	3275	29200	29223	53.7	41.7	5775	29358	29339	52.8	50	0.9	159	74.5	45.9	41	53.1
2256	3276	1046	1063	50.3		5776	1498	1481	51	50	0.7	453	76.4	43.9	41	53.7
2257	3277	24019	24039	50.1	42.9		24527	24508	50.5	45	0.4	509	75.4	41.1	41	52.9
2258	3278	24014	24035	50.6	40.9		24527	24508	50.5	45	0.1	514	75.5	41.2	41	53.1
2259	3279	1046	1063	50.3		5779	1497	1480	50.3	50	0.1	452	76.5	44	41	53.7
2260	3280	29201	29222	51	40.9		29358	29339	52.8	50	1.9	158	74.3	45.6	41	52.4
2261	3281	18583	18603	54.8	47.6		18696	18672	53.9	40	0.8	114	70.5	40.4	41	50.6
				تبب						.5	5.5	- 4	. 0.0	70.7		50.0

221	3282	12977	1299	50.	0 4	15700	1 40004	1 1 1 1 1 1								
220		23843				5782	13326				_			6 4	4 4	1 53.4
220		29200				10.00	24088								1 4	1 53.2
220		23843					29358						_			1 53
226		17792				5785	24091				_				4 4	1 53.3
226		23843				5786	18231						_		5 4	1 53.1
226		8374				5787	24094							45.	6 4	1 53.4
226		17793		_			9109			1		736	75.4	40.	1 4	1 53.1
	0 3290	1046				5789	18223				1.7	431	74.9	40.	4 4	1 52.5
227		23842				5790	1481	1463		1	0.2	436	76.2	43.	6 4	53.6
227						5791	24093				0	252	75.9	45.	6 - 4	
227		23842	23862			5792	24527	24507			0.1	686	76.1	41.	B 41	53.6
227		18550	2844			5793	3082	3058			1.9	260	74.3	41.	5 4	52.3
227		23841	18571	50.4			19316		50		0.4	767	75.5	40.	3 41	53
	6 3296		23860	52.1	55	10.00	24527	24507	51	42.9	1.1	687	76.1	41.	9 41	
227		23841	23860	52.1	55		24527	24508	50.5		1.6	687	76.1	41.9	41	
227		17793	17813	50			18233	18215	51.3	52.6	1.3	441	75.1	40.8	3 41	
227		23841	19		52.6		269	251	51.1	52.6	1.1	269	76.4	46.5	41	53.6
228		1	23859	50.5	-		24094	24076	50.9	52.6	0.4	254	76.1	46.1	41	53.5
228		8908	8925	51.1		5800	9249	9231	50.8	47.4	0.2	342	75.1	41.8	41	52.9
228		8908	8925	51.1		5801	9249	9230	51.5	45	0.5	342	75.1	41.8	41	53
		23841	23859	50.5		5802	24500	24481	50.1	45	0.4	660	76.1	42.1	41	53.4
228		23841	23859	50.5		5803	24526	24506	50.3	42.9	0.2	686	76.1	42	41	53.5
228		18225	18243	51.4		5804	18632	18611	50.2	40.9	1.2	408	75.7	42.6	41	53.2
		3794	3812	52.9		5805	4318	4294	54.4	40	1.5	525	75.5	41.1	41	53.8
228		8908	8925	51.1	50	5806	9245	9226	50	45	1	338	74.9	41.4	_	52.5
228		17790	17811	51.6	40.9	5807	18231	18210	52.2	45.5	0.6	442	75	40.5	41	53.1
228		18077	18100	54.7		5808	18443	18424	55.9	55	1.3	367	75.8	43.3	41	54.6
228		23838	23857	50.4		5809	24527	24507	51	42.9	0.6	690	76	41.7	41	53.4
	3310	23838	23857	50.4		5810	24527	24508	50.5	45	0.1	690	76	41.7	41	53.4
229		23735	23752	51.2		5811	24013	23995	50.3	47.4	0.8	279	74.1	40.5	41	52.1
2292		18080	18100	53.3	47.6		18220	18202	54.8	52.6	1.5	141	73.1	- 44	41	52.3
2293		18081	18100	51.7		5813	18223	18206	51.8	50	0.1	143	73.2	44.1	41	51.9
2294		18081	18100	51.7		5814	18231	18210	52.2	45.5	0.5	151	73.6	44.4	41	52.1
2295		18081	18100	51.7	50	5815	18233	18214	52	50	0.4	153	74	45.1	41	52.4
	3316	18081	18100	51.7		5816	18233	18215	51.3	52.6	0.4	153	74	45.1	41	52.3
2297	1	8911	8928	51.9		5817	9252	9235	50.1	50	1.8	342	75	41.5	41	52.6
2298		17791	17811	50	42.9		18223	18206	51.8	50	1.7	433	74.9	40.4	41	52.5
2299		8911	8928	51.9		5819	9249	9231	50.8	47.4	1	339	75	41.6	41	52.8
2300		12352	12375	52.9	41.7		12912	12892	53.6	52.4	0.8	561	76.2	42.6	41	54.3
2301	3321	8911	8928	51.9	50	5821	9249	9230	51.5	45	0.3	339	75	41.6	41	53
2302	3322	12352	12375	52.9		5822	12995	12976	51.1	45	1.8	644	76.4	42.9	41	53.9
2303		17791	17811	50	42.9	5823	18233	18215	51.3	52.6	1.3	443	75.1	40.9	41	52.7
2304		12977	12996	50.2	40	5824	13328	13307	51.2	45.5	1	352	76	44	41	53.4
2305		12977	12996	50.2	40	5825	13329	13308	50.5	40.9	0.3	353	76	43.9	41	53.4
2306		8911	8928	51.9	50	826	9245	9226	50	45	1.8	335	74.8	41.2	41	52.5
2307	3327	8913	8931	55.5	52.6	827	9252	9231	54	45.5	1.5	340	74.9	41.5	41	53.7
2308	3328	1402	1425	52.8	41.7	828	1501	1480	51.9	40.9	0.9	100	72	46	41	51.1
2309	3329	24941	24960	52	50 5	829	25646	25627	50.5	45	1.5	706	75.4	40.2	41	53
2310		17608	17628	50.9	42.9	830	17769	17749	50	42.9	0.9	162	72.4	40.7	41	50.8
2311	3331	24941	24960	52	50 5	831	25404	25386	52.7	52.6	0.7	464	75.3	41.2	41	53.4
2312	3332	24941	24960	52	50 5	832	25401	25383	50.6	47.4	1.4	461	75.2	41.2	41	53.4
									20.0	****		701	10.2	41	41	53

2313		24941	24960	52		5833	25400	25382	51.4	52.6	0.6	460	75.3	41.1	41	53.2
2314		17608	17628	50.9		5834	18231	18210	52.2	45.5	1.2	624	75.2	40.1	41	53.1
	3335	18081	18099	51.2	52.6		18232	18212	50.6	47.6	0.6	152	73.8	44.7	41	51.9
2316	3336	7725	7742	50		5836	7853	7833	50.7	47.6	0.7	129	71.2	40.3	41	49.9
2317	3337	8913	8931	55.5	52.6		9252	9230	54.5	43.5	1	340	74.9	41.5	41	53.9
2318	3338	17608	17628	50.9	42.9	5838	18233	18215	51.3	52.6	0.4	626	75.3	40.3	41	53.1
2319	3339	19715	19735	52.5	47.6	5839	19931	19912	50.9	55	1.6	. 217	74	41.9	41	52.2
2320	3340	8913	8931	55.5	52.6	5840	9248	9226	54.7	47.8	0.7	336	74.9	41.4	41	53.9
2321	3341	18081	18099	51.2	52.6	5841	18642	18622	50.5	42.9	0.7	562	76.2	42.7	41	53.6
2322	3342	2823	2844	50.4	45.5	5842	3189	3168	51	45.5	0.5	367	75.6	42.8	41	53.2
2323	3343	19715	19735	52.5	47.6	5843	19927	19908	52.1	55	0.3	213	73.9	41.8	41	52.4
2324	3344	2823	2844	50.4	45.5	5844	3190	3169	50.7	45.5	0.2	368	75.6	42.7	41	53.1
2325	3345	28936	28956	55.2	52.4	5845	29306	29287	54.6	55	0.6	371	76.6	45.3	41	55.1
2326	3346	28936	28956	55.2	52.4	5846	29306	29285	56.7	54.5	1.6	371	76.6	45.3	41	55.3
2327	3347	28523	28544	51.6	40.9	5847	29298	29280	51.4	52.6	0.2	776	78.4	47.3	41	55.4
2328	3348	24180	24199	50.3	40	5848	24933	24913	51.1	42.9	0.9	754	75.8	40.8	41	53.2
2329	3349	19715	19735	52.5	47.6	5849	19925	19905	51.4	52.4	1.1	211	73.8	41.7	41	52.2
2330	3350	4645	4665	50.2	42.9	5850	4836	4817	51.2	45	0.9	192	75	45.3	41	52.6
2331	3351	18081	18099	51.2	52.6	5851	18702	18685	50.2	50	1	622	76.2	42.4	41	53.5
2332	3352	28522	28542	50.2	42.9	5852	29298	29280	51.4	52.6	1.2	. 777	78.4	47.2	41	55.1
2333	3353	24179	24199	52.7	42,9	5853	24740	24717	52.5	41.7	0.2	562	76	42.2	41	54
2334	3354	19715	19735	52.5	47.6	5854	19909	19885	52.5	40	0	195	73.3	41	41	52.1
2335	3355	1810	1830	51.2	42.9	5855	2103	2082	52	45.5	0.8	294	75.5	43.5	41	53.3
2336	3356	19716	19737	52.2	45.5	5856	19909	19885	52,5	40	0.3	194	73.1	40.7	41	51.9
2337	3357	19719	19739	50.6	42.9	5857	19909	19885	52.5	40	1.9	. 191	72.9	40.3	41	51.3
2338	3358	12977	12996	50.2	40	5858	13332	13312	50.9	47.6	0.6	356	76.1	44.1	41	53.4
	3359	19721	19745	52.3	40	5859	19909	19885	52.5	40	0.2	189	73	40.7	41	51.9
2340	3360	17608	17627	50.2	45	5860	17769	17749	50	42.9	0.2	162	72.4	40.7	41	50.8
2341	3361	17608	17627	50.2	45	5861	18231	18210	52.2	45.5	2	624	75.2	40.1	41	52.8
2342		19794	19813	50	50	5862	20099	20078	50.5	40.9	0.5	306	74.4	40.8	41	52.2
2343		4658	4677	50.5	50	5863	5306	5289	50.8	50	0.3	649	75.5	40.7	41	53.1
2344		24179	24200	53.3	40.9	5864	24580	24560	51.3	52.4	2	402	74.6	40	41	52.7
2345		19794	19813	- 50	50	5865	19925	19906	50.1	50	0.1	132	72.8	43.9	41	51.1
2346	3366	1046	1064	51.2	47.4	5866	1498	1481	51	50	0.2	453	76.4	43.9	41	53.9
2347	3367	1046	1064	51.2	47.4	5867	1497	1480	50.3	50	0.9	452	76.5	44	41	53.7
2348		2133	2152	50.7	45		2675	2656	50.4	50	0.3	543	76.8	44.2	41	54
2349		28965	28984	52.9	55		29298	29279	52.6	55	0.3	334	76.4	45.2	41	54.4
2350		24378	24397	55		5870	24564	24542	55	47.8	0	187	73.6	42.2	41	53.1
2351		25348	25366	51.2	47.4		25651	25632	52.7	50	1.6	304	74.7	41.4	41	52.7
2352		19794	19813	50	50		19923	19903	50.9	47.6	0.9	130	72.7	43.8	41	51
2353		28967	28987	51.6	52.4		29358	29339	52.8	50	1.2	392	76.5	44.6	41	54.1
2354		17608	17627	50.2	45		18233	18215	51.3	52.6	1.1	626	75.3	40.3	41	52.9
2355		29186	29206	51.3		5875	29358	29339	52.8	50	1.5	173	74.5	45.1	41	52.6
2356		24379	24398	55		5876	25093	25074	54.6	55	0.4	715	75.9	41.4		54.6
2357		3170	3191	50.9	45.5		3646	3625	52	40.9	1.1	477	75.7	41.9	41	53.4
2358		3170	3191	50.9	45.5		3647	3628	50.6	45	0.3	478	75.7	42.1	41	53.3
2359		9140	9159	50.1	45		9249	9230	51.5	45	1.4	110	71.3	42.7	41	50
2360		12976	12995	51.1	45		13326	13306	50.7	42.9	0.4	351	76.1	44.2	41	53.6
2361		12976		51.1	45		13328	13307	51.2	45.5	0.1	353	76.1	44.2	41	53.7
2362		3168		51	45.5		3494	3473	50.4	40.9	0.5	327	75.1	42.2	41	52.8
2363		19794		50	50		19917	19896	50.9	45.5	0.9	124	72.3	43.5	41	50.7
2300	10000	10794	1 10013	1 30		10000	10017	10000		1 70.0	L 0.0	124	12.0	70.0	17'	1 00.1

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1		3384	24379	24398			5884	24517	24494	53.2		1.8	139	72.7	43.2	41	52
L	2365		24380	24399	55			25093	25074	54.6	55	0.4	714	76	41.5		54.6
L	2366		2823	2844	50.4			3201	3183	50.6	52.6	0.2	379	75.8	43	41	53.3
L	2367		3798	3819	54.2		1	4318	4294	54.4	40	0.2	521	75.4	41.1	41	54.2
L	2368	1000	9139	9159	52.5			9852	9829	53.1	45.8	0.6	714	75.4	40.1	41	53.6
L	2369	1	9139	9159	52.5	47.6	5889	9852	9828	53.6	44	1.1	714	75.4	40.1	41	53.6
L	2370	1	19795	19814	50.4	45	5890	19927	19908	52.1	55	1.7	133	72.7	43.6		51.1
L	2371		19795	19814	50.4	45	5891	19924	19905	50.1	50	0.3	130	72.4	43.1	41	50.8
L	2372		12976	12995	51.1	45	5892	13329	13308	50.5	40.9	0.6	354	76.1	44.1	41	53.5
. [2373		2133	2152	50.7	45	5893	2672	2654	50.9	52.6	0.2	540	76.8	44.3	41	54.1
	2374	1	4593	4613	51.5	47.6	5894	4836	4817	51.2	45	0.3	244	75.3	44.3	41	53.2
L	2375		1810	1830	51.2	42.9	5895	2113	2094	50.1	45	1.1	304	75.5	43.4	41	53
	2376	3396	17036	17058	53.5	47.8	5896	17483	17465	54.4	52.6	0.9	448	75.3	41.3	41	53.9
	2377		1046	1064	51.2	47.4	5897	1481	1463	50.5	47.4	0.6	436	76.2	43.6	41	53.6
L	2378		9055	9079	52.8	40	5898	9255	9236	51.1	45	1.8	201	73.5	41.3	41	51.9
L	2379		12976	12995	51.1	45	5899	13332	13312	50.9	47.6	0.2	357	76.2	44.3	41	53.7
L	2380	3400	8865	8884	50.4	45	5900	9311	9292	50.7	50	0.3	447	75.4	41.4	41	53
L	2381	3401	25363	25381	51.1	52.6	5901	25651	25632	52.7	50	1.6	289	74.3	40.8	41	52.5
	2382	3402	3168	3189	51	45.5	5902	3504	3485	50.4	45	0.6	337	75.3	42.4	41	52.9
E	2383	3403	25363	25381	51.1	52.6	5903	25649	25629	51.5	42.9	0.3	287	74.1	40:4	41	52.3
	2384	3404	29182	29202	51.2	42.9	5904	29414	29395	50.5	50	0.7	233	75.5	45.1	41	53.1
	2385	3405	3031	3051	51.3	52.4	5905	3497	3478	51.3	50	0.1	467	76.3	43.5	41	53.9
	2386	3406	29172	29192	51.5	42.9	5906	29412	29393	50.3	45	1.1	241	75.6	45.2	41	53.2
E	2387	3407	12040	12057	50.6	50	5907	12412	12392	50	42.9	0.6	373	75.9	43.4	41	53.2
	2388	3408	11543	11562	50.4	40	5908	12110	12090	51.1	42.9	0.7	568	75.9	41.9	41	53.4
	2389	3409	16909	16928	50.8	45	5909	17038	17021	50.7	50	0.1	130	72.7	43.8	41	51.2
	2390	3410	16909	16928	50.8	45	5910	17039	17022	51.4	50	0.6	131	72.6	43.5	41	51.2
	2391	3411	18077	18097	51.5	47.6	5911	18233	18214	52	50	0.5	157	73.9	44.6	41	52.3
	2392	3412	18077	18097	51.5	47.6	5912	18233	18215	51.3	52.6	0.2	157	73.9	44.6	41	52.2
	2393	3413	9055	9079	52.8	40	5913	9252	9234	51.4	52.6	1.4	198	73.7	41.9	41	52.1
L	2394	3414	25676	25697	51.9	40.9	5914	25832	25810	53.6	47.8	1.7	157	72.1	40:1	41	51.1
L	2395	3415	2223	2244	51.4	45.5	5915	2676	2657	50.7	50	0.7	454	76.9	45.2	41	54.2
L	2396	3416	619	640	50.4	45.5	5916	1171	1153	50.4	47.4	0	553	77.9	46.8	41	54.7
L	2397	3417	11541	11561	50.9	42.9	5917	12110	12090	51.1	42.9	0.3	570	75.9	41.9	41	53.5
L	2398		3360	3379	50.7	45	5918	3497	3478	51.3	50	0.6	138	74	46.4	42	52.1
L		3419	19725	19745	50	42.9	5919	19921	19901	50.2	47.6	0.1	197	73.5	41.6	42	51.6
_		3420	19720	19740	51.3	42.9	5920	19921	19901	50.2	47.6	1.1	202	73.4	41.1	42	51.5
	2401	3421	3360	3379	50.7	45	5921	3500	3481	51.2	50	0.5	141	74	46.1	42	52.1
		3422	19717	19738	50.8	40.9	5922	19921	19901	50.2	47.6	0.6	205	73.4	41	42	51.5
_		3423	24562	24580	50.1	52.6	5923	25209	25190	50.6	50	0.5	648	76.1	42	42	53.4
		3424	24559	24579	52	52.4	5924	24740	24717	52.5	41.7	0.5	182	76	48.4	42	53.9
_		3425	3360	3379	50.7	45	5925	3504	3485	50.4	45	0.3	145	74.2	46.2	42	52.2
_		3426	19716	19737	52.2	45.5	5926	19921	19900	51.8	45.5	0.4	206	73.5	41.3	42	52.1
		3427	3232	3251	50.3		5927	3494	3473	50.4	40.9	0.1	263	74.3	41.4	42	52.2
_		3428	26039	26058	54	55	5928	26657	26636	52.6	45.5	1.5	619	75.4	40.5	42	53.7
L-		3429	3232	3251	50.3	50	5929	3504	3485	50.4	45	0.1	273	74.6	41.8	42	52.4
		3430	19715	19735	52.5	47.6	5930	19921	19900	51.8	45.5	0.7	207	73.7	41.5	42	52.2
	2411		26039	26058	54	55	5931	26653	26631	53.2	43.5	0.9	615	75.3	40.3	42	53.8
		3432	3229	3248	50.6	50	5932	3494	3473	50.4	40.9	0.2	266	74.3	41.4	42	52.3
	2413		3229	3248	50.6	50	5933	3504	3485	50.4	45	0.3	276	74.5	41.7	42	52.4
L	2414	3434	3225	3244	52.4	55	5934	3495	3473	51.8	43.5	0.6	271	74.7	42.1	42	52.9
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2415	3435	3222	3241	52	50	5935	3650	3631	53.1	50	1.1	429	75.5	42	42	53.6
2416	3436	24559	24579	52	52.4	5936	25209	25190	50.6	50	1.3	651	76.1	42.1	42	53.6
2417	3437	6158	6178	51.3	42.9	5937	6289	6267	52.2	43.5	0.9	132	71.3	40.2	42	50.4
2418	3438	19709	19730	51.3	40.9		19917	19896	50.9	45.5	0.3	209	73.7	41.6	42	52
	3439	3223	3241	50.2	52.6		3497	3478	51.3	50	1.1	275	74.7	42.2	42	52.5
2420		3223	3241	50.2		5940	3646	3625	52	40.9	1.8	424	75.4	41.7	42	53
2421	3441	3223	3241	50.2		5941	3647	3628	50.6	45	0.4	425	75.5	41.9	42	53
2422		3217	3237	51.8		5942	3650	3631	53.1	50	1.3	434	75.5	41.9	42	53.5
2423		9352	9372	51.3	42.9	5943	10014	9996	50.7	52.6	0.6	663	75.6	40.7	42	53.2
2424		23733	23752	55.6		5944	24022	24003	55.5	55	0.1	290	74.5	41.4	42	53.9
2425		26040	26061	56.4		5945	26661	26639	55.3	47.8	1.2	622	75.5	40.7	42	54.5
	3446	9918	9938	51.4		5946	10608	10589	51	50	0.4	691	75.8	41.1	42	53.4
2427		7724	7742	51.4		5947	7843	7825	52.8	52.6	1.3	120	70.7	40	42	50
	3448	26040	26061	56.4		5948	26655	26631	56.2	48	0.2	616	75.4	40.6	42	54.8
	3449	28117	28135	50.6		5949	28506	28488	50.2	47.4	0.4	. 390	79.4	51.8	42	55.8
	3450	3217	3236	51.1		5950	3497	3478	51.3	50	0.2	281	74.7	42	42	52.7
2431		3217	3236	51.1	50	5951	3500	3481	51.2	50	0.1	284	74.7	41.9	42	52.7
2432		3165	3187	51.6		5952	3650	3631	53.1	50	1.5	486	75.8	42.2	42	53.6
	3453	19709	19730	51.3		5953	19925	19906	50.1	50	1.2	217	74	41.9	42	51.9
2434		9927	9945	50.8		5954	10199	10180	51.5	45	0.7	273	75.3	43.6	42	53.1
2435		9929	9946	50	50	5955	10670	10649	51.3	40.9	1.3	742	75.7	40.8	42	53.1
	3456	19709	19730	51.3		5956	19927	19908	52.1	55	0.9	219	74	42	42	52.3
2437	3457	9934	9953	50.7		5957	10356	10336	52.4	47.6	1.7	423	75.6	42.1	42	53.2
	3458	19709	19730	51.3		5958	19930	19910	50.6	47.6	0.7	222	74	41.9	42	52.1
2439		3164	3186	51.6		5959	3650	3631	53.1	50	1.5	487	75.9	42.3	42	53.7
	3460	3089	3110	51.8		5960	3188	3166	51.6	43.5	0.2	100	72	46	42	51
2441	3461	18979	19000	51.6		5961	19215	19194	50.2	40.9	1.4	237	73.5	40.1	42	51.6
2442		26421	26441	51.5		5962	26900	26882	51.5	52.6	0.1	480	77.5	46.2	42	54.8
2443		26421	26441	51.5		5963	26828	26810	52.9	52.6	1.4	408	76.6	44.9	42	54.2
2444		11540	11557	50.4		5964	11826	11802	51.3	40	0.8	287	74.4	41.1	42	52.3
2445		26421	26441	51.5		5965	26695	26678	50.5	50	1	275	74.9	42.5	42	52.7
	3466	11540	11557	50.4		5966	11819	11798	50.3	40.9	0.1	280	74.3	41.1	42	52.2
2447		11540	.11557	50.4		5967	11817	11797	50.4	42.9	0.1	278	74.3	41	42	52.2
2448	3468	23841	23859	50.5		5968	24515	24494	50.4	40.9	0.1	675	76.1	41.9	42	53.5
2449	3469	3055	3077	52.8		5969	3495	3473	51.8	43.5	0.9	441	76	43.1	42	53.9
2450	3470	3795	3813	52.1	52.6	5970	4565	4542	53.9	41.7	1.8	771	75.6	40.3	42	53.6
2451	3471	11540	11560	53.2		5971	11984	11966	53	52.6	0.2	445	75.1	40.7	42	53.6
2452	3472	11541	11561	50.9	42.9	5972	12165	12147	51.2	47.4	0.4	625	75.7	41.3	42	53.4
2453	3473	3795	3815	54.6	52.4	5973	4318	4294	54.4	40	0.2	524	75.5	41.2	42	54.3
2454	3474	7723	7741	52.2		5974	7853	7833	50.7	47.6	1.5	131	71.3	40.5	42	50.2
2455	3475	3055	3075	51.8	47.6	5975	3504	3485	50.4	45	1.4	450	76.1	43.1	42	53.5
2456	3476	3055	3074	51.1	50	5976	3494	3473	50.4	40.9	0.7	440	76	43	42	53.4
2457	3477	26421	26441	51.5		5977	26651	26631	50.2	42.9	1.3	231	73.8	41.1	42	51.8
2458	3478	28109	28130	50.2	40.9	5978	28672	28654	50.6	52.6	0.4	564	79.9	51.6	42	56.1
2459	3479	3055	3074	51.1	50	5979	3504	3485	50.4	45	0.7	450	76.1	43.1	42	53.5
2460	3480	12232	12250	51.9	52.6	5980	12412	12392	50	42.9	1.9	181	73.2	41.4	42	51.3
2461	3481	3034	3053	50.3	50	5981	3210	3190	50.5	47.6	0.2	177	74.9	45.8	42	52.6
2462	3482	3034	3053	50.3	50	5982	3494	3473	50.4	40.9	0.1	461	76.1	43.2	42	53.5
2463		3034	3053	50.3	50	5983	3504	3485	50.4	45	0.1	471	76.2	43.3	42	53.5
2464	3484	12236	12256	51.2	42.9	5984	12498	12480	50	47.4	1.1	263	74.6	42.2	42	52.4
2465	3485	12352	12375	52.9	41.7	5985	12724	12705	52.4	55	0.5	373	75.7	42.9	42	53.8

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2466 3		26421	26441	51.5	42.9	5986	26585	26567	51	47.4	0.5	165	72.2	40	42	50.9
2467 3		3031	3051	51.3	52.4	5987	3503	3484	51.5	50	0.1	473	76.3	43.6	42	53.9
2468 3		18704	18724	50.8	47.6	5988	19480	19459	50.3	40.9	0.4	777	75.5	40.2	42	53.1
2469 3		3016	3036	50.2	42.9	5989	3646	3625	52	40.9	1.8	631	76.4	42.8	42	53.6
2470 3		2823	2844	50.4	45.5	5990	3053	3034	50.3	50	0.2	231	74	41.6	42	52
2471 34		12366	12384	51.7	52.6	5991	12994	12976	50.3	47.4	1.3	629	76.4	42.9	42	53.7
2472 3		12366	12384	51.7	52.6	5992	12992	12974	51.2	52.6	0.5	627	76.5	43.1	42	54
2473 34		2823	2844	50.4	45.5	5993	3056	3037	52.1	55	1.6	234	74.2	41.9	42	52.2
2474 34		2522	2541	51.4	45	5994	2672	2654	50.9	52.6	0.5	151	75.3	48.3	42	53
2475 34		2522	2541	51.4		5995	2675	2656	50.4	50	1	154	75.2	48.1	_42	52.9
2476 34		2429	2447	50.2		5996	3056	3037	52.1	55	1.9	628	76.3	42.7	42	53.6
	497	2429	2447	50.2		5997	3190	3169	50.7	45.5	0.5	762	76.6	42.9	42	53.8
2478 34		27436	27455	52.7		5998	27541	27521	51.7	47.6	1	106	72.1	45.3	42	51.1
2479 34		2429	2447	50.2		5999	3192	3171	51.9	50	1.7	764	76.7	43.1	42	53.8
2480 35		2427	2445	52.1		6000	3056	3037	52.1	55	0	630	76.4	42.9	42	54.2
	501	27389	27407	50.6		6001	27541	27521	51.7	47.6	1.1	153	73.2	43.1	42	51.5
	502	2427	2445	52.1		6002	3190	3169	50.7	45.5	1.4	764	76.7	43.1	42	54
2483 35		18616	18636	51.4	47.6	6003	19316	19295	50	40.9	1.4	701	75.4	40.2	42	52.9
2484 35		2377	2395	52.4		6004	2672	2653	51.6	50	0.8	296	77	47.3	42	54.5
2485 35		18591	18611	51.7		6005	19216	19195	50.2	40.9	1.4	626	75.7	41.1	42	53.1
2486 35		12366	12384	51.7		6006	12739	12718	51	40.9	0.7	374	75.6	42.8	42	53.3
	507	2377	2395	52.4		6007	2672	2654	50.9	52.6	1.5	296	77	47.3	42	54.3
2488 35		16982	17001	51.2		6008	17111	17090	51.1	40.9	0.1	130	74.6	48.5	42	52.6
2489 35		2377	2395	52.4	52.6		2675	2656	50.4	50	2	299	77	47.2	42	54.1
2490 35		18590	18608	50.6	42.1		19216	19195	50.2	40.9	0.3	627	75.6	· 41	42	53.1
2491 35		2377	2395	52.4	52.6		2891	2873	50.8	47.4	1.6	515	76.8	44.5	42	54.1
2492 35		8220	8240	54	47.6		8935	8917	54.5	52.6	0.4	716	75.4	40.1	42	54.1
2493 35		12370	12388	50.1	47.4		12998	12979	50.1	45	0.1	629	76.4	42.9	42	53.6
2494 35		2223	2244	51.4	45.5		2675	2656	50.4	50	1	453	77	45.3	42	54.1
2495 35		2220	2239	51.3		6015	2672	2654	50.9	52.6	0.4	453	77	45.3	42	54.2
2496 35		24418	24439	52.9	45.5		24936	24919	51.8	50	1.1	519	76	42.4	42	53.8
2497 35		18586	18603	50.4	44.4		19216	19195	50.2	40.9	0.2	631	75.6	40.9	42	53.1
2498 35		2220	2239	51.3		6018	2675	2656	50.4	50	0.8	456	76.9	45.2	42	54.1
2499 35		1402	1422	50.2	42.9		2153	2134	50.4	45	0.2	752	76.7	43.1	42	53.8
2500 35	20	1356	1375	53.8	55	6020	2153	2133	52.1	42.9	1.7	798	76.9	43.5	42	54.5

Table 5: Primers

ID NO &	d primer SEQ Co-ordinates	Revers ID NO 8	e primer SEQ & Co-ordinates	T _M (FOR 8	REV) (°C)	Product length (bp)
6076	1-19	6171	199-183	50.1	50.3	199
6077	149-169	6172	334-315	51.5	52.4	186
6078	292-310	6173	560-541	50.8	51.1	269
6079	598-619	6174	749-731	52.6	50.6	152
6080	721-742	6175	930-912	50.4	50.3	210
6081	888-912	6176	1077-1058	52.8	51.2	190
6082	984-1003	6177	1149-1131	51.1	51.1	166

6083	1157-1175	6178	1479-1460	50.9	51.6	323
6084	1420-1441	6179	1700-1680	51.2	50.7	281
6085	1685-1707	6180	1834-1811	53.8	53.7	150
6086	1740-1764	6181	1987-1963	53.4	52.2	248
6087	2007-2025	6182	2251-2232	50.3	50.1	245
6088	2226-2245	6183	2385-2366	50.4	50.1	160
6089	2428-2446	6184	2749-2728	50.1	50.3	322 .
6090	2742-2763	6185	2893-2875	50.6	51.4	152
6091	2823-2844	6186	3082-3058	50.4	52.3	260
6092	3007-3031	6187	3185-3164	51.9	51.0	179
6093	3234-3254	6188	3497-3478	51.1	51.3	264
6094	3453-3476	6189	3647-3627	51.8	52.1	195
6095	3601-3622	6190	3877-3853	52.5	53.6	277
6096	4007-4027	6191	4158-4135	51.1	51.4	. 152
6097	4141-4165	6192	4316-4295	51.3	50.8	176
6098	4366-4387	6193	4567-4544	54.6	55.4	202
6099	· 4488-4508	6194	4708-4690	50.7	50.3	221
6100	4658-4677	6195	4994-4974	50.5	51.2	337
6101	4902-4922	6196	5115-5092	50.5	51.4	214
6102	5239-5260	6197	5450-5430	50.8	50.9	212
6103	5366-5389	6198	5560-5542	50.5	51.8	195
6104	5593-5612	6199	5860-5836	50.8	51.6	. 268
6105	6042-6062	6200	6291-6271	50.4	51.1	250
6106	6271-6291	6201	6483-6463	51.1	50.2	213
6107	7017-7040	6202	7171-7153	52.4	52.8	155
6108	7253-7272	6203	7504-7486	50.3	50.3	252
6109	7415-7434	6204	. 7677-7654	54.5	53.6	263
6110	7615-7635	6205	7821-7798	51.1	52.8	207
6111	7728-7746	6206	7936-7915	51.7	50.1	209
6112	7845-7867	6207	7994-7970	52.7	53.4	150
6113	8011-8029	6208	8189-8170	51.4	50.6	179
6114	8143-8166	6209	8300-8281	52.2	50.8	158
6115	8221-8239	6210	8388-8369	51.0	51.1	168
6116	8553-8575	6211	8931-8915	51.8	50.3	379
6117	8867-8886	6212	9254-9236	50.7	50.6	388

6118	9244-9267	6213	9597-9573	51.9	53.4	354
6119	9620-9640	6214	9990-9969	51.3	51.3	371
6120	10009-10027	6215	10188-10171	50.2	50.2	180
6121	10093-10113	6216	10244-10223	52.4	50.6	152
6122	10242-10265	6217	10608-10589	51.2	51.0	367
6123	10549-10571	6218	10783-10763	53.7	55.2	235
6124	10766-10785	6219	10930-10912	52.0	51.1	165 -
6125	11065-11085	6220	11305-11287	50.7	50.0	241
6126	11265-11287	6221	11429-11405	54.5	53.5	165
6127	11552-11571	6222	11730-11709	52.0	50.4	179
6128	11705-11726	6223	11869-11848	50.1	50.2	165
6129	11801-11824	6224	11984-11967	51.5	50.4	184
6130	12040-12058	6225	12254-12235	52.3	51.9	215
6131	12235-12253	6226	12406-12388	50.1:	50.1	· 172
6132	12366-12384	6227	12730-12712	51.7	52.2	365
6133	12727-12748	6228	12994-12976	50.8	50.3	268
6134	12948-12966	6229	13224-13201	50.7	51.7	277
6135	, 13175-13196	6230	13324-13300	54.3	55.1	150
. 6136	13237-13258	6231	13545-13526	52.9	52.9	309
6137	13790-13810	6232	13963-13945	50.9	50.7	174
6138	14080-14098	6233	14280-14257	51.5	51.0	201
-6139	14405-14427	6234	14561-14540	50.2	50.9	157
6140	14882-14906	6235	15046-15024	50.9	51.5	165
6141	14951-14976	6236	15145-15124	53.1	52.9	195
6142	15113-15134	6237	15275-15257	51.6	50.8	163
6143	15211-15230	6238	15383-15363	50.2	50.1	173
6144	15364-15387	6239	15528-15506	54.0	52.1	165
6145	15456-15477	6240	15605-15585	52.0	53.2	150
6146	15513-15532	6241	15897-15876	51.2	50.4	385
6147	15837-15856	6242	15999-15978	52.3	50.8	163
6148	16073-16096	6243	16301-16277	51.7	52.8	229
6149	16245-16266	6244	16404-16380	50.3	52.0	160
6150	16366-16385	6245	16515-16492	52.9	53.8	150
6151	16553-16571	6246	16777-16758	53.4	51.5	225
6152	16832-16852	6247	17026-17004	51.0	51.6	195

6153	16982-17001	6248	17359-17340	51.2	50.2	378
6154	17354-17372	6249	17511-17490	51.3	50.4	158
6155	17422-17443	6250	17573-17552	50.2	51.1	152
6156	17603-17623	6251	17769-17748	50.7	51.5	167
6157	17728-17746	6252	17883-17862	50.9	51.2	156
6158	18011-18030	6253	18163-18140	52.9	51.9	153
6159	18076-18098	6254	18225-18205	54.4	55.0	150 .
6160	18270-18292	6255	18432-18413	51.9	51.4	163
6161	18352-18373	6256	18648-18629	51.3	50.8	297
6162	18550-18571	6257	18702-18684	50.4	51.9	153
6163	18720-18738	6258	19004-18983	50.6	51.0	285
6164	18960-18981	6259	19109-19085	54.7	54.3	150
6165	19065-19089	6260	19217-19195	52.8	51.7	153
6166	19310-19329	6261	19476-19454	50.2	52.1	167
. 6167	19569-19589	6262	19719-19701	50.5	51.8	151
6168	.19707-19731	6263	19856-19833	55.7	55.9	150
6169	19771-19792	6264	19921-19901	50.1	50.2	151
6170	19833-19851	6265	19986-19966	50.9	50.7	154

Table 6: Primers

	rd primer SEQ & Co-ordinates		se primer SEQ & Co-ordinates	T _M (FOR &	REV) (°C)	Product length (bp)
6266	20110-20132	6305	20425-20404	51.9	50.9	316
6267	20468-20492	6306	20617-20596	53.2	53.5	150
6268	20557-20578	6307	20891-20871	50.4	50.6	335
6269	20838-20856	6308	21037-21015	52.5	52.0	200
6270	21096-21116	6309	21295-21272	50.1	51.7	200
6271	22173-22194	6310	22414-22395	52.4	51.0	242
6272	22320-22342	6311	22501-22479	54.8	54.3	182
6273	22532-22552	6312	22695-22675	50.6	50.0	164
6274	22712-22736	6313	22873-22852	56.7	55.5	162
6275	22842-22861	6314	23086-23067	51.0	52.8	245
6276	23151-23170	6315	23395-23376	51.4	50.3	245
6277	23307-23326	6316	23524-23501	51.1	51.1	218
6278	23615-23635	6317	23776-23758	50.7	50.2	162

6279	23838-23857	6318	23996-23977	50.4	50.6	159
6280	24030-24051	6319	24407-24386	57.6	55.7	378
6281	24388-24407	6320	24581-24563	50.4	50.1	194
6282	24559-24579	6321	24938-24921	52.0	50.4	380
6283	24922-24941	6322	25184-25166	50.1	51.2	263
6284	25201-25220	6323	25400-25382	51.1	51.4	200
6285	25363-25381	6324	25646-25627	51.1	50.5	284 -
6286	25656-25681	6325	25839-25814	54.5	56.4	184
6287	25761-25782	6326	25982-25961	54.6	54.3	222
6288	26039-26058	6327	26189-26166	54.0	53.0	151
6289	26184-26205	6328	26333-26310	50.9	51.8	150
6290	26422-26442	6329	26660-26641	51.3	50.2	239
6291	26571-26589	6330	26739-26715	51.7	53.2	169
6292	26733-26752	6331	26960-26941	51.1	52.2	· 228
6293	26866-26885	6332	27139-27117	50.7	51.9	274
6294	27300-27321	6333	27458-27439	51.2	50.2	. 159
6295	27361-27380	6334	27579-27558	52.4	51.1	219
6296	27718-27740	6335	27917-27901	50.7	50.0	200
6297	28041-28059	6336	28207-28189	50.8	50.8	167
6298	28166-28189	6337	28411-28393	52.2	52.9	246
6299	28395-28414	6338	28671-28653	51.5	50.2	277
6300	28654-28672	6339	28821-28800	50.6	52.3	168
6301	28867-28885	6340	29184-29166	51.5	51.6	318
6302	29183-29204	6341	29360-29342	50.4	50.4	178
6303	29262-29279	6342	29626-29606	50.1	50.2	365
6304	29538-29559	6343	29690-29670	50.0	50.4	153

Table 7: Primers

Name	SEQ ID NO:	Co-ordinates	Name	SEQ ID NO:	Co-ordinates
AB4f	6344	19869-19888	CB1r	6367	28011-28030
AB5f	6345	20238-20257	CB2r	6368	27671-27690
BC1f	6346	20581-20600	CB3r	6369	27301-27320
BC2f	6347	20950-20969	CB4r	6370	26931-26950
BC3f	6348	21339-21358	CB5r	6371	26575-26594
BC4f	6349	21708-21727	CB6r	6372	26191-26210
BC5f	6350	22041-22060	CB7r	6373	25841-25860
BC6f	6351	22410-22429	CB8r	6374	25476-25495
BC7f	6352	22759-22778	CB9r	6375	25126-25145

BC9f 6354 23500-23519 CB11r 6377 24422-24 BC10f 6355 23841-23860 CB12r 6378 24031-24 BC11f 6356 24210-24229 CB13r 6379 23673-23 BC12f 6357 24560-24579 CB14r 6380 23298-23 BC13f 6358 24941-24960 CB15r 6381 22928-22 BC14f 6359 25310-25329 CB16r 6382 22567-22 BC15f 6360 25675-25694 CB17r 6383 22196-22						
BC10f 6355 23841-23860 CB12r 6378 24031-24 BC11f 6356 24210-24229 CB13r 6379 23673-23 BC12f 6357 24560-24579 CB14r 6380 23298-23 BC13f 6358 24941-24960 CB15r 6381 22928-22 BC14f 6359 25310-25329 CB16r 6382 22567-22 BC15f 6360 25675-25694 CB17r 6383 22196-22	BC8f	6353	23131-23150	CB10r	6376	24791-24810
BC11f 6356 24210-24229 CB13r 6379 23673-23 BC12f 6357 24560-24579 CB14r 6380 23298-23 BC13f 6358 24941-24960 CB15r 6381 22928-23 BC14f 6359 25310-25329 CB16r 6382 22567-22 BC15f 6360 25675-25694 CB17r 6383 22196-22	BC9f	6354	23500-23519	CB11r	6377	24422-24441
BC12f 6357 24560-24579 CB14r 6380 23298-23 BC13f 6358 24941-24960 CB15r 6381 22928-23 BC14f 6359 25810-25329 CB16r 6382 22567-22 BC15f 6360 25675-25694 CB17r 6383 22196-22	BC10f	6355	23841-23860	CB12r	6378	24031-24050
BC13f 6358 24941-24960 CB15r 6381 22928-22 BC14f 6359 25310-25329 CB16r 6382 22567-22 BC15f 6360 25675-25694 CB17r 6383 22196-22	BC11f	6356	24210-24229	CB13r	6379	23673-23692
BC14f 6359 25310-25329 CB16r 6382 22567-22 BC15f 6360 25675-25694 CB17r 6383 22196-22	BC12f	6357	24560-24579	CB14r	6380	23298-23317
BC15f 6360 25675-25694 CB17r 6383 22196-22	BC13f	6358	24941-24960	CB15r	6381	22928-22947
	BC14f	6359	25310-25329	CB16r	6382	22567-22586
PC16f coct occor CP10- cont otopt of	BC15f	6360	25675-25694	CB17r	6383	22196-22215
BC101 6361 20044-20063 CB181 6384 21831-21	BC16f	6361	26044-26063	CB18r	6384	21831-21850
BC17f 6362 26413-26432 CB19r 6385 21431-21	BC17f	6362	26413-26432	CB19r	6385	21431-21450
BC18f 6363 26763-26782 CB20r 6386 21073-21	BC18f	6363	26763-26782	CB20r	6386	21073-21092
BC19f 6364 27132-27151 CB21r 6387 20715-20	BC19f	6364	27132-27151	CB21r	6387	20715-20734
BC20f 6365 27491-27510 BA1r 6388 20345-20	BC20f	6365	27491-27510	BA1r	6388	20345-20364
BC21f 6366 27845-27864 BA2r 6389 19969-19	BC21f	6366	27845-27864	BA2r	6389	19969-19988
BA3r 6390 19599-19				BA3r	6390	19599-19618
BA4r 6391 19228-19				BA4r	6391	19228-19247
BA5r 6392 18852-18		L		BA5r	6392	18852-18871

Table 8: Primers

Name	SEQ ID NO	Co-ordinates	Name	SEQ ID NO	Co-ordinates
. F1	6393	1-19	- R1	6441	334-315
F2	6394	292-310	R2	6442	749-731
F3	6395	721-742	R3	6443	1077-1058
F4	6396	984-1003	- R4	6444	1479-1460
F5	6397	1420-1441	· R5	6445	1834-1811
F6	6398	1740-1764	R6	6446	2251-2232
F7	6399	2226-2245	. R7	6447	2749-2728
F8	6400	2742-2763	: R8	6448	3082-3058
F9	6401	3007-3031	: R9	6449	3497-3478
F10	6402	3453-3476	· R10	6450	3877-3853
F11	6403	4007-4027	R11	6451	4316-4295
F12	6404	4366-4387	R12	6452	4708-4690
F13	6405	4658-4677	R13	6453	5115-5092
F14	6406	5239-5260	R14	6454	5560-5542
F15	6407	5593-5612	R15	6455	6291-6271
F16	6408	6271-6291	R16	6456	7171-7153
F17	6409	7253-7272	R17	6457	7677-7654
F18	6410	7615-7635	R18	6458	7936-7915
F19	6411	7845-7867	R19	6459	8189-8170
F20	6412	8143-8166	R20	6460	8388-8369
F21	6413	8553-8575	R21	6461	9254-9236
F22	6414	9244-9267	R22	6462	9990-9969
F23	6415	10009-10027	R23	6463	10244-10223
F24	6416	10242-10265	R24	6464	10783-10763
F25	6417	10766-10785	R25	6465	11305-11287
F26	6418	11265-11287	R26	6466	11730-11709
F27	6419	11705-11726	R27	6467	11984-11967
F28	6420	12040-12058	R28	6468	12406-12388
F29	6421	12366-12384	R29	6469	12994-12976
F30	6422	12948-12966	R30	6470	13324-13300
F31	6423	13237-13258	R31	6471	13963-13945

F32	6424	14080-14098	R32	6472	14561-14540
F33	6425	14882-14906	R33	6473	15145-15124
F34	6426	15113-15134	R34	6474	15383-15363
F35	6427	15364-15387	R35	6475	15605-15585
F36	6428	15513-15532	R36	6476	15999-15978
F37	6429	16073-16096	R37	6477	16404-16380
F38	6430	16366-16385	R38	6478	16777-16758
F39	6431	16832-16852	R39	6479	17359-17340
F40	6432	17354-17372	R40	6480	17573-17552
F41	6433	17603-17623	R41	6481	17883-17862
F42	6434	18011-18030	R42	6482	18225-18205
F43	6435	18270-18292	R43	6483	18648-18629
F44	6436	18550-18571	R44	6484	19004-18983
F45	6437	18960-18981	R45	6485	19217-19195
F46	6438	19310-19329	R46	6486	19719-19701
F47	6439	19707-19731	R47	6487	19921-19901
F48	6440	19833-19851		0.	.002.110001

Table 9: Primers

	Name	SEQ ID NO:
1	CB12R	6488
2	. R0010	6489
3	R0011	6490
4	R0012	6491
5	BNI-ED	6492
6	BNI-EU	6493
7	SAR1S-U	6494
8	SAR1As-D	6495
9	SAR1S	6496
10	SAR1As	6497
11	IN2-U	6498
12	IN4-D	6499
13	IN-2	6500
14	IN-4	6501
15	IN-6	6502
16	IN-7	6503
17	COR1-U	6504
18	COR2-D	6505
19	COR-1	6506
20	COR-2	6507
21	HKUF-U	6508
22	HKUR-D	6509
23	HKU-F	6510
24	HKU-R	6511
25	1451-D	6512
26	1451-U	6513
27	690-D	6514
28	690-U	6515
29	690-D2	6516

	Name	SEQ ID NO:
37	EMC8-D2	6524
38	EMC8-U2	6525
39	EMC11-D	6526
40	EMC11-U	6527
41	ORF1B-D	6528
42	ORF1B-U	6529
43	ORFS-D	6530
44	ORFS-U	6531
45	E7-717F	6532
46	E8-85R	6533
47	E8-307F	6534
48	E11-771F	6535
49	E11-96R	6536
50	CON1-F	6537
_51	CON1-U	6538
52	CON2-F	6539 .
53	CON2-R	6540
54	CON3-F	6541
55	CON3-R	6542
_56	15-F	6543
57	15-R	6544
58	15-F2	6545
59	15-R2	6546
60	13-F	6547
61	13-R	6548
62	13-F2	6549
_63	13-R2	6550
64	CONTIG-F	6551
65	QT3-R	6552

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	30	690-U2	6517
	31	EMC7-D	6518
	32	EMC7-U	6519
ļ	33	EMC7-D2	6520
	34	EMC7-U2	6521
	35	EMC8-D	6522
	36	EMC8-U	6523

66	QT3-F	6553
67	QIN-R	6554
68	QIN-F	6555
69	AB1-F	6556
70	AB2-F	6557
71	AB3-F	6558
72	AB1-R	6559

Table 10: Features of the predicted proteins and open reading frames of the SARS virus

	SARS ORF	Length	Role	Cleavage	Features	Cons
	(SEQ ID NO)	(aa)		site		*
	P28 (9766)	179	Leader protein	179 (G/G)*		+
	P65 (9767)	639	Homologue of MHV p65 cleavage product	818 (G/A)		+
	Nsp1 (9768)	2422 **	Papain like protease, cleaves the first two proteins	3240 (Q/S)	phosphoesterase domain Zn bindlng domain	+
ORF1a	Nsp2 (9769)	306	3C-Ilke protease, cleaves proteins nsp1-nsp12	3546 (Q/G)		+.
	Nsp3 (9770)	290	?	3836 (Q/S)	5 TMDs	+
	Nsp4 (9771)	83	?	3919 (Q/A)	1 TMD	+
	Nsp5 (9772)	198	?	4117 (Q/N)		+
	Nsp6 (9773)	113	?	4230 (Q/A)		+
	Nsp7 (9774)	139	?	4369 (Q/S)	Putative growth factor-like motif	+
	Nsp9 (9775)	932	RNA polymerase	5298 (Q/A)		+
ORF1b	Nsp10 (9776)	601	Putative helicase Tenner et al (2003) J Biol Chem 278:39578-82	5899 (Q/A)	Metal binding domain, ATP/GTP binding domain	+
8	Nsp11 (9777)	527	?	6426 (Q/S)	2 January derman	+ 1
	Nsp12 (9778)	346	?	6772 (Q/A)		+
	Nsp13 (9779)	298	?	-		+
	Spike (S) (6042)		Major antigenic determinant, contains the receptor-binding domain		Leader peptide, 1 TMD, 17 N- glycosylation sites	+
	Orf3 (6043)	274	?		2 TMDs, 1 N-glycosylation site, 10 O-glycosylation sites	-
	Orf4 (6044)	154	?			-
_	Envelope (E) (6045)	76	Associated with viral envelope		1 TMD, 2 N-glycosylation sites	+
Structural region	Matrix (M) (6046)	221	Associated with viral envelope, membrane spanning protein	OO!	3 TMDs, 1 N-glycosylation site	+ ·
Struct	Orf7 (6047)	63	?		1 TMD	-
ļ	Orf8 (6048)	122	?		1 TMD	
-	Orf9 (6049)	44	?		Surface-associated	
ļ	Orf10	39	?		Surface-associated	
-	Orf11(6050)	84	?		1 N-glycosylation site	-
	Nucleocapsid (N) (6052)	422	Associated with viral genomic RNA		phosphoprotein	+
	Orf13	98	?		1 O-glycosylation site	

TMD: predicted transmembrane domain.

Cons^d *: + indicates presence of corresponding protein at least in one of the other coronaviruses #: Alternatively, cleaved after Gly-Gly (i.e. at G/A) to give a 180mer

##: This 2422mer may be further cleaved after residue 1922 (Gly-2740 of SEQ ID NO: 6039) to give a 1922mer PLpro containing the Zn-binding motif (SEQ ID NO: 7254) and a 500mer.

Table 11: Protein homologies between SARS and other coronaviruses

Numbers indicate percentage of aminoacid identity between SARS proteins and corresponding gene products of other coronaviruses. More conserved pairs are in bold; more variable pairs are underlined.

		group 1		gro	up 2	group 3
Proteins	229E	TGV	PEDV	MHV	BC ₀ V	AIBV
REPLICASE REGION						
leader protein p28	<20	<20	<20	27	<20	<20
p65 homologue	<20	23	23	<20	20	<20
nsp1 (PLP protease)	25.5	25.8	25.4	29	30	<u>25</u>
nsp2 (3CL protease)	40.4	43.8	44.6	50	48.4	41
nsp3	30	<u>27</u>	29.4	34.2	35.5	28.5
nsp4	38.6	42.2	39.8	47.5	46.1	37.3
nsp5	48.2	42.9	43.9	46.8	47.3	38.7
nsp6	45.1	38.9	45.1	45.1	46.9	39.8
nsp7	<u>53.8</u>	54.5	56.1	56.2	55.4	58.3
nsp9 (polymerase)	59.8	59.6	60	67.3	66.9	62.4
nsp10 (helicase)	60.7	62	62.3	67.2	68.6	<u>58.9</u>
nsp11	52.3	53.7	52.3	57.6	57.6	<u>52</u>
nsp12	43.1	43	45.4	45.9	45	40,2
nsp13 .	56.4	54.4	55,3	63	65	<u>53.4</u>
STRUCTURAL REGION			•			
Spike (S)	28.8	31.6*	30.3	31.1	31	32.7*
Envelope (E)	33*	27.9	20	23	26.5	23.2
Matrix glycoprotein (M)	<u>30.6</u>	32.5	34.8	40.8	41.9	32.5
Nucleocapsid (N)	26.9	30.1	29.5	37.3	37.4	31.5

^{*} These three alignments were obtained only on a fragment of the whole protein.

5

Table 12: Nucleotide and aminoacid differences between five SARS isolates

		FRA*	TOR2*	Urbani*	CUHK*	HKU*
	position°	base/aminoacid	base/aminoacid	base/aminoacid	base/aminoacid	base/aminoacio
	2557	A/Thr	G/Ala	G/Ala		
	2601	T/Val	WAIA	G/Ala	G/Ala	G/Ala
	7746	G/Pro				С
	7919	C/Ala		T/Val	Т	
	7930	G/Asp		1/Vai		
ORF1a	8387	G/Ser				A/Asn
	8416	G/Arg				C/Thr
	9404	T/Val				C/Thr
	9479	T/Val			C/Ala	
	11448	T/lle	; C		C/Ala	
		17116		С	С	СС
	13494	GT/Val				AG/Ser
	16622	C/Ala		т .		Adroei
	17564	T/Asp			C/Glu	
ORF1b	17846	C/Arg			T	
OIN ID	18065	G/Lys				A
	18965	A/IIe	T	T	- -	
	19064	A/Glu		G	G	
	19084	T/IIe	C/Thr	C/Thr	C/Thr	C/Thr
	21721	G/Glv			0/0	
	22222	T/IIe			A/Asp C/Thr	
spike	23220	T/Ser	G/Ala		C/Thr	
	24872	T/Leu	G// UL	С		
	24933	T/Phe	C/Leu	C/Leu	C/Leu	C/Leu
	25298	G/Gly	A/Arg			0,200
ORF3	25569	T/Met	Avaig			A.0.
						A/Lys
matrix	26600	T/Val	C/Ala	C/Ala	C/Ala	
	26857	T/Ser		C/Pro		
ORF10	27827	T/Cys			C/Arg	
nucleocapsid	28268	T/Ile	C/Thr	C/Thr	C/Thr	C/Thr

^{*} SARS coronavirus FRA (accession number AY310120)

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SARS coronavirus TOR2 (accession number AY274119)

SARS coronavirus Urbani (accession number AY278741) SARS coronavirus CUHK-W1 (accession number AY278554)

SARS coronavirus HKU-39849 (accession number AY278491)

[°] The position is based on the FRA sequence.

5

Epitope predictions were performed at http://www.mpiib-berlin.mpg.de/MAPPP/binding.html using a minimum score of 0.5 and the BIMAS matrix, with a maximum of 20 results being selected. The analysis revealed 9mer and 10mer epitopes.

Table 13: Epitopes for SEQ ID NO: 6039

-	HLA A1 - 9 mers						
Maximu	Maximum possible score using this molecule type 5625						
Rank	Start position	Sequence	% of max. score	Score			
1	1867	SEQ ID NO: 7400	8 %	450			
2	4139	SEQ ID NO: 7401	5.55 %	312.5			
3	88	SEQ ID NO: 7402	4 %	225			
4 ·	4249	SEQ ID NO: 7403	3.55 %	200			
5	4059	SEQ ID NO: 7404	2.22 %	125			
6	2027	SEQ ID NO: 7405	1.6 %	90			
7	3413	SEQ ID NO: 7406	1.11 %	62.5			
8	1823	SEQ ID NO: 7407	0.88 %	50			
9	2798	SEQ ID NO: 7408	0.88 %	50			
10	220	SEQ ID NO: 7409	0.8 %	45			
11	3738	SEQ ID NO: 7410	0.8 %	45			
12	4182	SEQ ID NO: 7411	0.8 %	45			
13	4174	SEQ ID NO: 7412	0.66 %	37.5			
14	1940	SEQ ID NO: 7413	0.55 %	31.25			
15	38	SEQ ID NO: 7414	0.48 %	27			
16	1231	SEQ ID NO: 7415	0.44 %	25			
17	1613	SEQ ID NO: 7416	0.44 %	25			
18	3645	SEQ ID NO: 7417	0.44 %	25			
19	4192	SEQ ID NO: 7418	0.44 %	25			
20	378	SEQ ID NO: 7419	0.4 %	22.5			

	HLA A1 - 10 mers						
Maximu	m possible score us	sing this molecule type		5625			
Rank Start position Sequence % of max. score							
1	1867	SEQ ID NO: 7420	8 %	450			
2	1495	SEQ ID NO: 7421	4 %	225			
3	3921	SEQ ID NO: 7422	2.4 %	135			
4	486	SEQ ID NO: 7423	2.22 %	125			
5	4139	SEQ ID NO: 7424	2.22 %	125			

6	62	SEQ ID NO: 7425	1.6 %	90
7	1190	SEQ ID NO: 7426	1.6 %	90
8	1284	SEQ ID NO: 7427	1.6 %	90
9	3284	SEQ ID NO: 7428	1.6 %	90
10	2921	SEQ ID NO: 7429	1.2 %	67.5
11	349	SEQ ID NO: 7430	0.8 %	45
12	789	SEQ ID NO: 7431	0.8 %	45
13	1185	SEQ ID NO: 7432	0.8 %	45
14	4184	SEQ ID NO: 7433	0.8 %	45
15	1313	SEQ ID NO: 7434	0.64 %	36
16	3948	SEQ ID NO: 7435	0.48 %	27
17	149	SEQ ID NO: 7436	0.44 %	25
18	941	SEQ ID NO: 7437	0.44 %	25
19	1390	SEQ ID NO: 7438	0.44 %	25
20	1613	SEQ ID NO: 7439	0.44 %	25

HLA A3 - 9 mers						
Maximum possible score using this molecule type						
Rank	Start position	Sequence	% of max. score	Score		
1	1010	SEQ ID NO: 7440	1.48 %	180		
2	3155	SEQ ID NO: 7441	1.48 %	180		
3	1229	SEQ ID NO: 7442	1.23 %	150		
4	2405	SEQ ID NO: 7443	0.88 %	108		
5	2	SEQ ID NO: 7444	0.74 %	90		
6	2304	SEQ ID NO: 7445	0.74 %	90		
7	2358	SEQ ID NO: 7446	0.74 %	90		
8	3160	SEQ ID NO: 7447	0.74 %	90		
.9	3771	SEQ ID NO: 7448	0.74 %	90		
10	4007	SEQ ID NO: 7449	0.74 %	90		
11	3079	SEQ ID NO: 7450	0.66 %	81		
12	4045	SEQ ID NO: 7451	0.66 %	81		
13	1081	SEQ ID NO: 7452	0.49 %	60		
14	3268	SEQ ID NO: 7453	0.49 %	60		
15	4144	SEQ ID NO: 7454	0.49 %	60		
16	614	SEQ ID NO: 7455	0.37 %	45		
17	728	SEQ ID NO: 7456	0.37 %	45		
18	1537	SEQ ID NO: 7457	0.37 %	45		
19	313	SEQ ID NO: 7458	0.32 %	40		
20	1744	SEQ ID NO: 7459	0.32 %	40		

	HLA A3 - 10 mers					
Maximu	ım possible score us	sing this molecule type		12150		
Rank	Start position	Sequence	% of max. score	Score		
1	62	SEQ ID NO: 7460	4.44 %	540		
2	2151	SEQ ID NO: 7461	2.46 %	300		
3	633	SEQ ID NO: 7462	2.22 %	270		
4	1158	SEQ ID NO: 7463	2.22 %	-270		
5	2565	SEQ ID NO: 7464	2.22 %	270		
6	2298	SEQ ID NO: 7465	1.77 %	216		
7	3159	SEQ ID NO: 7466	1.11 %	135		
8	640	SEQ ID NO: 7467	0.98 %	120		
9	2186	SEQ ID NO: 7468	0.74 %	90		
10	3869	SEQ ID NO: 7469	0.74 %	90		
11	2308	SEQ ID NO: 7470	0.66 %	81		
12	786	SEQ ID NO: 7471	0.55 %	67.5		
13	749	SEQ ID NO: 7472	0.49 %	60		
14	1080	SEQ ID NO: 7473	0.49 %	60		
15	2358	SEQ ID NO: 7474	0.49 %	60		
16	3955	SEQ ID NO: 7475	0.49 %	60		
17	714	SEQ ID NO: 7476	0.37 %	45		
18	1081	SEQ ID NO: 7477	0.37 %	45		
19	1170	SEQ ID NO: 7478	0.37 %	45		
20	1228	SEQ ID NO: 7479	0.37 %	45		

	HLA A24 - 9 mers					
Maximu	ım possible score ι	using this molecule ty	ре	1596.672		
Rank	Start position	Sequence	% of max. score	Score		
1	3797	SEQ ID NO: 7480	37.57 %	600		
2	4202	SEQ ID NO: 7481	37.57 %	600		
3	3189	SEQ ID NO: 7482	25.05 %	400		
4	1864	SEQ ID NO: 7483	23.14 %	369.6		
5	1066	SEQ ID NO: 7484	22.54 %	360		
6	2143	SEQ ID NO: 7485	22.54 %	360		
7	2693	SEQ ID NO: 7486	22.54 %	360		
8	1426	SEQ ID NO: 7487	18.78 %	300		
9	1238	SEQ ID NO: 7488	18.03 %	288		
10	3768	SEQ ID NO: 7489	18.03 %	288		
11	797	SEQ ID NO: 7490	15.03 %	240		

12	1882	SEQ ID NO: 7491	15.03 %	240		
13	1490	SEQ ID NO: 7492	13.77 %	220		
_14	2237	SEQ ID NO: 7493	13.77 %	220		
15	95	SEQ ID NO: 7494	12.52 %	200		
16	1821	SEQ ID NO: 7495	12.52 %	200		
17	2289	SEQ ID NO: 7496	12.52 %	200		
18	3080	SEQ ID NO: 7497	12.52 %			
19	3660	SEQ ID NO: 7498	12.52 %	200		
20	4354	SEQ ID NO: 7499	12.52 %	200		
	200 12.52 % 200					

<u></u>	HLA A24 - 10 mers						
Maximu	Maximum possible score using this molecule type 1596.672						
	Start position	Sequence	% of max. score	Score			
11	2143	SEQ ID NO: 7500	37.87 %	604.8			
2	1159	SEQ ID NO: 7501	26.30 %	420			
3	1650	SEQ ID NO: 7502	26.30 %	420			
4	1150	SEQ ID NO: 7503	18.78 %	300			
5	2763	SEQ ID NO: 7504	18.78 %	. 300			
6	3165	SEQ ID NO: 7505	18.78 %	300			
7	3201	SEQ ID NO: 7506	15.03 %	240			
8	3694	SEQ ID NO: 7507	15.03 %	240			
9	4204	SEQ ID NO: 7508	15.03 %	240			
10	1692	SEQ ID NO: 7509	13.77 %	220			
11	797	SEQ ID NO: 7510	12.52 %	200			
12	1610	SEQ ID NO: 7511	12.52 %	200			
13	1789	SEQ ID NO: 7512	12.52 %	200			
14	1881	SEQ ID NO: 7513	12.52 %	200			
15	3090	SEQ ID NO: 7514	12.52 %	200			
16	3763	SEQ ID NO: 7515	12.52 %	200			
17	2569	SEQ ID NO: 7516	11.27 %	180			
18	194	SEQ ID NO: 7517	9.39 %	150			
19	1771	SEQ ID NO: 7518	9.39 %	150			
20	2488	SEQ ID NO: 7519	9.39 %	150			

HLA A 0201 - 9 mers					
Maximum possible score using this molecule type 3925227.1					
Rank	Start position		% of max. score		
1		SEQ ID NO: 7520		8144.13515256	
_ 2	3729	SEQ ID NO: 7521	0.10 %	4047.23088	

3	3574	SEQ ID NO: 7522	0.09 %	3547.4996634
4	3615	SEQ ID NO: 7523	0.06 %	2722.682592
5	3159	SEQ ID NO: 7524	0.05 %	1999.734264
6	2339	SEQ ID NO: 7525	0.03 %	1551.92907744
7	2201	SEQ ID NO: 7526	0.03 %	1521.53694
8	3559	SEQ ID NO: 7527	0.02 %	1174.38939504
9	3085	SEQ ID NO: 7528	0.02 %	1146.296448
10	4070	SEQ ID NO: 7529	0.02 %	970.4103696
11	3708	SEQ ID NO: 7530	0.02 %	958.92888
12	3098	SEQ ID NO: 7531	0.02 %	942.678
13	1362	SEQ ID NO: 7532	0.02 %	900.6984
14	3563	SEQ ID NO: 7533	0.01 %	735.86016
15	3774	SEQ ID NO: 7534	0.01 %	687.655656
16	4242	SEQ ID NO: 7535	0.01 %	685.78272
17	2340	SEQ ID NO: 7536	0.01 %	668.37342936
18	650	SEQ ID NO: 7537	0.01 %	640.1983392
19	3862	SEQ ID NO: 7538	0.01 %	620.57772
20 ·	2860	SEQ ID NO: 7539	0.01 %	607.88448

	HLA A 0201 - 10 mers			
	Maximum possible score using this molecule type			3925227.1
Rank	Start position	Sequence	% of max. score	Score
1	2307	SEQ ID NO: 7540	0.40 %	15915.66281448
2	`2201	SEQ ID NO: 7541	0.12 %	4772.09313
3	3558	SEQ ID NO: 7542	0.05 %	2295.04855632
4	1772	SEQ ID NO: 7543	0.04 %	1759.6656
5	3087	SEQ ID NO: 7544	0.03 %	1215.76896
6	2339	SEQ ID NO: 7545	0.02 %	1116.29986272
7	2308	SEQ ID NO: 7546	0.02 %	970.14776112
8	3061	SEQ ID NO: 7547	0.02 %	836.2525104
9	2748	SEQ ID NO: 7548	0.01 %	726.706344
10	3837	SEQ ID NO: 7549	0.01 %	720.8292
11	59	SEQ ID NO: 7550	0.01 %	650.3112
12	2877	SEQ ID NO: 7551	0.01 %	620.22996
13	4114	SEQ ID NO: 7552	0.01 %	559.8936
14	805	SEQ ID NO: 7553	0.01 %	484.4565072
15	1655	SEQ ID NO: 7554	0.01 %	437.48208
_16	611	SEQ ID NO: 7555	0.00 %	319.9392
17	1961	SEQ ID NO: 7556	0.00 %	305.94186

18	1223	SEQ ID NO: 7557	0.00 %	289.08792
19	852	SEQ ID NO: 7558	0.00 %	285.67242
20	2139	SEQ ID NO: 7559	0.00 %	284.845869

HLA A 1101 - 9 mers				
Maximu	ım possible score u	sing this molecule type		36
Rank	Start position	Sequence	% of max. score	Score
11	4200	SEQ ID NO: 7560	50 %	18
2	281	SEQ ID NO: 7561	25 %	9
3	3236	SEQ ID NO: 7562	25 %	9
4	509	SEQ ID NO: 7563	16.66 %	6
5	848	SEQ ID NO: 7564	16.66 %	6
6	2193	SEQ ID NO: 7565	16.66 %	6
7	3542	SEQ ID NO: 7566	16.66 %	6
8	541	SEQ ID NO: 7567	15 %	5.4
9	1748	SEQ ID NO: 7568	12.5 %	4.5
10	829	SEQ ID NO: 7569	11.11 %	4
11	1149	SEQ ID NO: 7570	11.11 %	4
12	2027	SEQ ID NO: 7571	11.11 %	4
13	2576	SEQ ID NO: 7572	11.11 %	4
14	873	SEQ ID NO: 7573	8.33 %	3
15	2725	SEQ ID NO: 7574	8.33 %	3
16	3541	SEQ ID NO: 7575	8.33 %	3
17	. 1837	SEQ ID NO: 7576	7.5 %	2.7
18	2475	SEQ ID NO: 7577	7.5 %	2.7
19	2703	SEQ ID NO: 7578	7.5 %	2.7
20	1823	SEQ ID NO: 7579	6.66 %	2.4

HLA A 1101 - 10 mers				
		sing this molecule type		36
Rank	Start position	Sequence	% of max. score	Score
11	3541	SEQ ID NO: 7580	50 %	18
2	281	SEQ ID NO: 7581	25 %	9
3	1495	SEQ ID NO: 7582	25 %	9
4	2303	SEQ ID NO: 7583	25 %	9
5	2616	SEQ ID NO: 7584	25 %	9
6	48	SEQ ID NO: 7585	16.66 %	6
7	1394	SEQ ID NO: 7586	16.66 %	6
8	1499	SEQ ID NO: 7587	16.66 %	6

9	1862	SEQ ID NO: 7588	16.66 %	6
10	1163	SEQ ID NO: 7589	11.11 %	4
11	4006	SEQ ID NO: 7590	11.11 %	4
12	4344	SEQ ID NO: 7591	11.11 %	4
13	633	SEQ ID NO: 7592	10 %	3.6
14	119	SEQ ID NO: 7593	8.33 %	3
15	1190	SEQ ID NO: 7594	8.33 %	3
16	1195	SEQ ID NO: 7595	8.33 %	- 3
17	1725	SEQ ID NO: 7596	8.33 %	3
.18	2728	SEQ ID NO: 7597	8.33 %	3
19	2895	SEQ ID NO: 7598	8.33 %	3
20	3033	SEQ ID NO: 7599	8.33 %	3

HLA B7 - 9 mers				
Maximu	m possible score us	sing this molecule type		5400
Rank	Start position	Sequence	% of max. score	Score
1	1335	SEQ ID NO: 7600	4.44 %	240
2	2580	SEQ ID NO: 7601	4.44 %	240
3	1703	SEQ ID NO: 7602	3.70 %	200
4	113	SEQ ID NO: 7603	2.22 %	120
5	168	SEQ ID NO: 7604	2.22 %	120
6	2842	SEQ ID NO: 7605	2.22 %	120
7	4027	SEQ ID NO: 7606	2.22 %	120
8	3680	SEQ ID NO: 7607	1.66 %	90
9	2085	SEQ ID NO: 7608	1.48 %	80
10	2492	SEQ ID NO: 7609	1.48 %	80
11	2660	SEQ ID NO: 7610	1.48 %	80
12	2906	SEQ ID NO: 7611	1.48 %	80
13	3346	SEQ ID NO: 7612	1.48 %	80
14	4038	SEQ ID NO: 7613	1.48 %	80
15	1163	SEQ ID NO: 7614	1.11 %	60
16	1457	SEQ ID NO: 7615	1.11 %	60
17	2351	SEQ ID NO: 7616	1.11 %	60
18	2471	SEQ ID NO: 7617	1.11 %	60
19	3499	SEQ ID NO: 7618	1.11 %	60
20	3635	SEQ ID NO: 7619	1.11 %	60

HLA B7 - 10 mers	
Maximum possible score using this molecule type	5400

Rank	Start position	Sequence	% of max. score	Score
11	1703	SEQ ID NO: 7620	3.70 %	200
_ 2	17	SEQ ID NO: 7621	2.22 %	120
3	3008	SEQ ID NO: 7622	2.22 %	120
4	4106	SEQ ID NO: 7623	2.22 %	120
5	3450	SEQ ID NO: 7624	1.66 %	90
6	. 113	SEQ ID NO: 7625	1.48 %	80
7	195	SEQ ID NO: 7626	1.48 %	- 80
8	307	SEQ ID NO: 7627	1.48 %	- 80
9	780	SEQ ID NO: 7628	1.48 %	80
10.	1000	SEQ ID NO: 7629	1.48 %	80
11	1072	SEQ ID NO: 7630	1.48 %	80
12	1404	SEQ ID NO: 7631	1.48 %	80
13	1980	SEQ ID NO: 7632	1.48 %	80
14	2262	SEQ ID NO: 7633	1.48 %	80
15	2543	SEQ ID NO: 7634	1.48 %	80
16	2906	SEQ ID NO: 7635	1.48 %	80
17	3077 ·	SEQ ID NO: 7636	1.48 %	80
18	3175	SEQ ID NO: 7637	1.48 %	. 80
19	4195	SEQ ID NO: 7638	1.48 %	80
20	4251	SEQ ID NO: 7639	1.48 %	80

Table 14: Epitopes for SEQ ID NO: 6040

HLA A1 - 9 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	20	SEQ ID NO: 7640	0.04 %	2.25	
2	91	SEQ ID NO: 7641	0.01 %	1	
3	125	SEQ ID NO: 7642	0.01 %	0.75	
4	56	SEQ ID NO: 7643	0.00 %	0.5	
5	145	SEQ ID NO: 7644	0.00 %	0.5	

	HLA A1 - 10 mers				
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	20	SEQ ID NO: 7645	0.01 %	0.9	
2	56	SEQ ID NO: 7646	0.00 %	0.5	
3	71	SEQ ID NO: 7647	0.00 %	0.5	

4	144	SEQ ID NO: 7648	0.00 %	0.5

	HLA A3 - 9 mers				
Maximu	m possible score us	sing this molecule type		12150	
Rank	Start position	Sequence	% of max. score	Score	
1	115	SEQ ID NO: 7649	· 0.24 %	30	
2	87	SEQ ID NO: 7650	0.04 %	6	
3	80	SEQ ID NO: 7651	0.03 %	4.05	
4	125	SEQ ID NO: 7652	0.01 %	1.8	
5	39	SEQ ID NO: 7653	0.01 %	1.5	
6	56	SEQ ID NO: 7654	0.01 %	1.5	
7	135	SEQ ID NO: 7655	0.00 %	1.2	
- 8	91	SEQ ID NO: 7656	0.00 %	1	
9	119	SEQ ID NO: 7657	0.00 %	1	
10	141	SEQ ID NO: 7658	0.00 %	0.9	
11	150	SEQ ID NO: 7659	0.00 %	0.6	
12	137	SEQ ID NO: 7660	0.00 %	0.54	

HLA A3 - 10 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	36	SEQ ID NO: 7661	0.24 %	30	
2	144	SEQ ID NO: 7662	0.06 %	8	
3	101	SEQ ID NO: 7663	0.03 %	4	
4	99	SEQ ID NO: 7664	0.02 %	3.6	
5	80	SEQ ID NO: 7665	0.02 %	2.7	
6	125	SEQ ID NO: 7666	0.01 %	1.6875	
. 7	71	SEQ ID NO: 7667	0.01 %	1.5	
8	118	SEQ ID NO: 7668	0.01 %	1.5	
9	40	SEQ ID NO: 7669	0.01 %	1.35	
10	5	SEQ ID NO: 7670	0.00 %	0.9	
11	56	SEQ ID NO: 7671	0.00 %	0.9	
12	107	SEQ ID NO: 7672	0.00 %	0.6	
13	135	SEQ ID NO: 7673	0.00 %	0.6	
14	141	SEQ ID NO: 7674	0.00 %	0.6	
15	148	SEQ ID NO: 7675	0.00 %	0.6	
16	116	SEQ ID NO: 7676	0.00 %	0.5	

HLA A24 - 9 mers

Maximu	Maximum possible score using this molecule type			1596.672
Rank	Start position	Sequence	% of max. score	Score
1	153	SEQ ID NO: 7677	1.05 %	16.8
2	80	SEQ ID NO: 7678	0.75 %	12
3	123	SEQ ID NO: 7679	0.50 %	8
4	137	SEQ ID NO: 7680	0.50 %	8
5	9	SEQ ID NO: 7681	0.45 %	7.2
6	77	SEQ ID NO: 7682	0.45 %	7.2
7	112	SEQ ID NO: 7683	0.45 %	7.2
8	73	SEQ ID NO: 7684	0.41 %	6.6
9	32	SEQ ID NO: 7685	0.37 %	6
10	110	SEQ ID NO: 7686	0.37 %	6
11	140	SEQ ID NO: 7687	0.37 %	6
12	143	SEQ ID NO: 7688	0.37 %	6
13	. 18	SEQ ID NO: 7689	0.30 %	4.8
14	54	SEQ ID NO: 7690	0.30 %	4.8
15	108	SEQ ID NO: 7691	0.30 %	4.8
16	141	SEQ ID NO: 7692	0.30 %	4.8
17	92	SEQ ID NO: 7693	0.27 %	4.4
18	33	SEQ ID NO: 7694	0.25 %	4
19	49	SEQ ID NO: 7695	0.25 %	4
20	111	SEQ ID NO: 7696	0.25 %	4

	HLA A24 - 10 mers				
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
11	142	SEQ ID NO: 7697	12.52 %	200	
2	110	SEQ ID NO: 7698	0.75 %	.12	
3	99	SEQ ID NO: 7699	0.50 %	8	
4	8	SEQ ID NO: 7700	0.45 %	7.2	
5	140	SEQ ID NO: 7701	0.45 %	7.2	
6	32	SEQ ID NO: 7702	0.37 %	6	
7	17	SEQ ID NO: 7703	0.30 %	4.8	
8	53	SEQ ID NO: 7704	0.30 %	4.8	
9	76	SEQ ID NO: 7705	0.30 %	4.8	
10	107	SEQ ID NO: 7706	0.30 %	4.8	
11	111	SEQ ID NO: 7707	0.30 %	4.8	
12	72	SEQ ID NO: 7708	0.27 %	4.4	
13	91	SEQ ID NO: 7709	0.27 %	4.4	

14	31	SEQ ID NO: 7710	0.25 %	4
15	127	SEQ ID NO: 7711	0.25 %	4
16	139	SEQ ID NO: 7712	0.25 %	4
17	80	SEQ ID NO: 7713	0.22 %	3.6
18	38	SEQ ID NO: 7714	0.18 %	3
19	118	SEQ ID NO: 7715	0.18 %	3
20	49	SEQ ID'NO: 7716	0.12 %	2

	HLA A 0201 - 9 mers				
Maxim	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	80	SEQ ID NO: 7717	0.00 %	171.96732	
2	147	SEQ ID NO: 7718	0.00 %	51.46848	
3	143	SEQ ID NO: 7719	0.00 %	11.6146182	
4	56	SEQ ID NO: 7720	0.00 %	11.304684	
5	10	SEQ ID NO: 7721	0.00 %	10.34586	
6	6	SEQ ID NO: 7722	0.00 %	6.56830734	
7	26	SEQ ID NO: 7723	0.00 %	6.07614	
8	141	SEQ ID NO: 7724	0.00 %	5.981472	
9	148	SEQ ID NO: 7725	0.00 %	5.194044	
10	9	SEQ ID NO: 7726	0.00 %	4.299183	
11	137	SEQ ID NO: 7727	0.00 %	4.299183	
12	130	SEQ ID NO: 7728	0.00 %	4.138344	
13	84	SEQ ID NO: 7729	0.00 %	3.42792	
14	27	SEQ ID NO: 7730	0.00 %	3.383484	
15	2	SEQ ID NO: 7731	0.00 %	3.381	
16	62	SEQ ID NO: 7732	0.00 %	3.251556	
17	23	SEQ ID NO: 7733	0.00 %	2.9542005	
18	99	SEQ ID NO: 7734	0.00 %	1.982232	
19	33	SEQ ID NO: 7735	0.00 %	1.86921	
20	111	SEQ ID NO: 7736	0.00 %	1.76402985	

	HLA A 0201 - 10 mers				
Maxim	um possible score	using this molecule	type	3925227.1	
Rank	Rank Start position Sequence % of max. score				
1	5	SEQ ID NO: 7737	0.00 %	159.9696	
2	25	SEQ ID NO: 7738	0.00 %	69.552	
3	80	SEQ ID NO: 7739	0.00 %	36.5148	
4	107	SEQ ID NO: 7740	0.00 %	21.3624	

5	148	SEQ ID NO: 7741	0.00 %	17.73576
6	61	SEQ ID NO: 7742	0.00 %	13.9104
7	147	SEQ ID NO: 7743	0.00 %	11.304684
88	53	SEQ ID NO: 7744	0.00 %	8.230458
9	17	SEQ ID NO: 7745	0.00 %	7.3086111
10	110	SEQ ID NO: 7746	0.00 %	6.174104475
11	9	SEQ ID NO: 7747	0.00 %	6.0858
12	99	SEQ ID NO: 7748	0.00 %	5.6823984
13	2 .	SEQ ID NO: 7749	0.00 %	3.188283
14	41	SEQ ID NO: 7750	0.00 %	2.206413
15	135	SEQ ID NO: 7751	0.00 %	2.076624
16	76	SEQ ID NO: 7752	0.00 %	2.005692
17	23	SEQ ID NO: 7753	0.00 %	1.798209
18	40	SEQ ID NO: 7754	0.00 %	1.68996456
19	39	SEQ ID NO: 7755	0.00 %	1.516482
20	118	SEQ ID NO: 7756	0.00 %	1.2683304

HLA A 1101 - 9 mers				
Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score
1	91	SEQ ID NO: 7757	2.77 %	1

	HLA A 1101 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	101	SEQ ID NO: 7758	33.33 %	12	
2	71	SEQ ID NO: 7759	2.77 %	1	
3	90	SEQ ID NO: 7760	1.66 %	0.6	

	HLA B7 - 9 mers				
Maximu	m possible score us	sing this molecule type		5400	
Rank	Start position	Sequence	% of max. score	Score	
1	49	SEQ ID NO: 7761	2.22 %	120	
2	9	SEQ ID NO: 7762	1.11 %	60	
3	73	SEQ ID NO: 7763	0.66 %	36	
4	33	SEQ ID NO: 7764	0.37 %	20	
5	137	SEQ ID NO: 7765	0.37 %	20	
6	141	SEQ ID NO: 7766	0.37 %	20	
7	77	SEQ ID NO: 7767	0.22 %	12	

8	112	SEQ ID NO: 7768	0.22 %	12
9	143	SEQ ID NO: 7769	0.22 %	12
10	81	SEQ ID NO: 7770	0.14 %	8
11	13	SEQ ID NO: 7771	0.09 %	5
12	69	SEQ ID NO: 7772	0.09 %	5
13	18	SEQ ID NO: 7773	0.07 %	4
14	32	SEQ ID NO: 7774	0.07 %	4
15	54	SEQ ID NO: 7775	0.07 %	- 4
16	80	SEQ ID NO: 7776	0.07 %	4
17	92	SEQ ID NO: 7777	0.07 %	4
18	108	SEQ ID NO: 7778	0.07 %	4
19	111	SEQ ID NO: 7779	0.07 %	· 4
20	123	SEQ ID NO: 7780	0.07 %	4

HLA B7 - 10 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	99	SEQ ID NO: 7781	0.74 %	40	
2	17	SEQ ID NO: 7782 0:37 %		20	
3	8	SEQ ID NO: 7783	0.22 %	. 12	
4	72	SEQ ID NO: 7784	0.22 %	12	
5	91	SEQ ID NO: 7785	0.22 %	. 12	
6	127	SEQ ID NO: 7786	0.11 %	6	
7	31	SEQ ID NO: 7787	0.07 %	4	
8	32	SEQ ID NO: 7788 0.07 %		· 4	
9	53:	SEQ ID NO: 7789 0.07 %		4	
10	76 .	SEQ ID NO: 7790 0.07 %		4	
11	107	SEQ ID NO: 7791	0.07 %	4	
12	110	SEQ ID NO: 7792	0.07 %	4	
13	111	SEQ ID NO: 7793	0.07 %	4	
14	140	SEQ ID NO: 7794	0.07 %	4	
15	9	SEQ ID NO: 7795	0.05 %	3	
16	19	SEQ ID NO: 7796	0.05 %	3	
17	33	SEQ ID NO: 7797 0.03 %		2	
18	93	SEQ ID NO: 7798 0.03 %		2	
19	102	SEQ ID NO: 7799	0.03 %	2	
20	129	SEQ ID NO: 7800	0.02 %	1.5	

Table 15: Epitopes for SEQ ID NO: 6041

HLA A1 - 9 mers				
Maximu	m possible score u	sing this molecule type		5625
Rank	Start position	Sequence	% of max. score	Score
11	1818	SEQ ID NO: 7801	1.6 %	90
2	373	SEQ ID NO: 7802 1.33 %		75
3	681	SEQ ID NO: 7803	1.33 %	75
4	74	SEQ ID NO: 7804	0.88 %	50
5	786	SEQ ID NO: 7805	0.88 %	50
6	1495	SEQ ID NO: 7806	0.88 %	50
_ 7	88	SEQ ID NO: 7807	0.8 %	45
8	. 357	SEQ ID NO: 7808	0.8 %	45
9	1271	SEQ ID NO: 7809	0.8 %	45
10	1799	SEQ ID NO: 7810	0.8 %	45
11	1393	SEQ ID NO: 7811	0.48 %	• 27
12	386	SEQ ID NO: 7812	0.44 %	- 25
13	. 2304	SEQ ID NO: 7813	0.44 %	. 25
14	198	SEQ ID NO: 7814	0.4 %	22.5
15	840	SEQ ID NO: 7815	0.4 %	22.5
16	2359	SEQ ID NO: 7816	0.4 %	22.5
17	1194	SEQ ID NO: 7817	0.32 %	18
18	1546	SEQ ID NO: 7818	0.32 %	18
19	2200	SEQ ID NO: 7819	0.22 %	12.5
20	996	SEQ ID NO: 7820	0.2 %	11.25

HLA A1 - 10 mers					
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	995	SEQ ID NO: 7821	10 %	562.5	
2	1303	SEQ ID NO: 7822	2.22 %	125	
3	1582	SEQ ID NO: 7823	2 %	112.5	
4	1456	SEQ ID NO: 7824	1.6 %	90	
5	772	SEQ ID NO: 7825	1.11 %	62.5	
6	181	SEQ ID NO: 7826	0.88 %	50	
7	632	SEQ ID NO: 7827	0.88 %	50	
8	2281	SEQ ID NO: 7828	0.88 %	50	
9	1586	SEQ ID NO: 7829	0.8 %	45	
10	2109	SEQ ID NO: 7830	0.8 %	45	
11	745	SEQ ID NO: 7831	0.55 %	31.25	

1916	SEQ ID NO: 7832	0.53 %	30
966	SEQ ID NO: 7833	0.44 %	25
1387	SEQ ID NO: 7834	0.44 %	25
2263	SEQ ID NO: 7835	0.44 %	25
2457	SEQ ID NO: 7836	0.26 %	15
1057	SEQ ID NO: 7837		12.5
2562	SEQ ID NO: 7838		12.5
74	SEQ ID NO: 7839		10
298	SEQ ID NO: 7840		10
	966 1387 2263 2457 1057 2562 74	966 SEQ ID NO: 7833 1387 SEQ ID NO: 7834 2263 SEQ ID NO: 7835 2457 SEQ ID NO: 7836 1057 SEQ ID NO: 7837 2562 SEQ ID NO: 7838 74 SEQ ID NO: 7839	966 SEQ ID NO: 7833 0.44 % 1387 SEQ ID NO: 7834 0.44 % 2263 SEQ ID NO: 7835 0.44 % 2457 SEQ ID NO: 7836 0.26 % 1057 SEQ ID NO: 7837 0.22 % 2562 SEQ ID NO: 7838 0.22 % 74 SEQ ID NO: 7839 0.17 %

HLA A3 - 9 mers				
Maximu		sing this molecule type	9	12150
Rank	Start position	Sequence	% of max. score	Score
1	536	SEQ ID NO: 7841	3.33 %	405
2	986	SEQ ID NO: 7842	2.46 %	300
3	805	SEQ ID NO: 7843	1.64 %	200
4	2345	SEQ ID NO: 7844	1.48 %	180
5	2481	SEQ ID NO: 7845	0.55 %	67.5
6	204	SEQ ID NO: 7846	0.49 %	60
_ 7	895	SEQ ID NO: 7847	0.44 %	54
8	1512	SEQ ID NO: 7848	0.44 %	54
9	2491	SEQ ID NO: 7849	0.37 %	45
10	436	SEQ ID NO: 7850	0.32 %	40
11	917	SEQ ID NO: 7851	0.32 %	40
12	1176	SEQ ID NO: 7852	0.32 %	40
13	1517	SEQ ID NO: 7853	0.29 %	36
14	466	SEQ ID NO: 7854	0.24 %	30
15	1784	SEQ ID NO: 7855	0.24 %	30
16	2039	SEQ ID NO: 7856	0.24 %	30
17	2124	SEQ ID NO: 7857	0.24 %	30
18	1049	SEQ ID NO: 7858	0.22 %	27
19	2200	SEQ ID NO: 7859	0.22 %	27
20	2598	SEQ ID NO: 7860	0.22 %	27

	HLA A3 - 10 mers				
Maximum possible score using this molecule type 12					
Rank	Start position	Sequence	% of max. score	Score	
1	392	SEQ ID NO: 7861	2.46 %	300	
2	2230	SEQ ID NO: 7862	1.48 %	180	

3	590	SEQ ID NO: 7863	1.11 %	135
4	697	SEQ ID NO: 7864	1.11 %	135
5	919	SEQ ID NO: 7865	0.74 %	90
6	1354	SEQ ID NO: 7866	0.74 %	90
7	1430	SEQ ID NO: 7867	0.74 %	90
8	2534	SEQ ID NO: 7868	0.74 %	90
9	202	SEQ ID NO: 7869	0.49 %	60
10	488	SEQ ID NO: 7870	0.49 %	60
11	922	SEQ ID NO: 7871	0.49 %	60
12	1735	SEQ ID NO: 7872	0.49 %	60
13	2281	SEQ ID NO: 7873	0.49 %	- 60
14	1894	SEQ ID NO: 7874	0.44 %	54
15	2552	SEQ ID NO: 7875	0.44 %	54
16	555	SEQ ID NO: 7876	0.37 %	. 45
17	1134	SEQ ID NO: 7877	0.37 %	· 45
18	1149	SEQ ID NO: 7878	0.29 % :	36
_19	283	SEQ ID NO: 7879	0.24 %	30
20	917	SEQ ID NO: 7880	0.24 %	30
	317	2EQ ID NO: 7880	0.24 %	30

HLA A24 - 9 mers					
Maximu	Maximum possible score using this molecule type 1596.672				
Rank	Start position	Sequence	% of max. score	Score	
1	2375	SEQ ID NO: 7881	36.07 %	576	
2	1751	SEQ ID NO: 7882	28.93 %	462	
3	195	SEQ ID NO: 7883	25.05 %	400	
4	2306	SEQ ID NO: 7884	21.04 %	336	
5	806	SEQ ID NO: 7885	20.66 %	330	
6	1252	SEQ ID NO: 7886	18.78 %	300	
7	160	SEQ ID NO: 7887	15.03 %	240	
8	517	SEQ ID NO: 7888	15.03 %	240	
9	375	SEQ ID NO: 7889	12.52 %	200	
10	1275	SEQ ID NO: 7890	12.52 %	200	
11	2175	SEQ ID NO: 7891	12.52 %	200	
12	2207	SEQ ID NO: 7892	12.52 %	200	
13	2343	SEQ ID NO: 7893	12.52 %	200	
_14	443	SEQ ID NO: 7894	11.27 %	180	
15	668	SEQ ID NO: 7895	7.51 %	120	
16	1825	SEQ ID NO: 7896	6.88 %	110	
17	1690	SEQ ID NO: 7897	4.69 %	75	

18	159	SEQ ID NO: 7898	3.75 %	60
19	2550	SEQ ID NO: 7899	3.75 %	60
20	1949	SEQ ID NO: 7900	3.38 %	54

HLA A24 - 10 mers				
Maximu	ım possible score ı	using this molecule ty	pe	1596.672
Rank	Start position	Sequence	% of max. score	Score
1	641	SEQ ID NO: 7901	45.09 %	720
_2	809	SEQ ID NO: 7902	24.80 %	396
3	1209	SEQ ID NO: 7903	22.54 %	360
4	216	SEQ ID NO: 7904	18.03 %	288
5	159	SEQ ID NO: 7905	15.03 %	240
6	528	SEQ ID NO: 7906	15.03 %	240
7	799	SEQ ID NO: 7907	15.03 %	240
8	1436	SEQ ID NO: 7908	15.03 %	240
9	2219	SEQ ID NO: 7909	15.03 %	240
10	1065	SEQ ID NO: 7910	13.77 %	220
11	1953	SEQ ID NO: 7911	13.15 %	210
12	1966	SEQ ID NO: 7912	12.52 %	200
13	2600	SEQ ID NO: 7913	12.52 %	200
14	71	SEQ ID NO: 7914	9.39 %	150
15	380	SEQ ID NO: 7915	9.39 %	150
16	1989	SEQ ID NO: 7916	9.39 %	150
17	342	SEQ ID NO: 7917	8.76 %	140
18	1071	SEQ ID NO: 7918	8.76 %	140
19	2570	SEQ ID NO: 7919	6.88 %	110
20	2550	SEQ ID NO: 7920	6.26 %	100

	HLA A 0201 - 9 mers				
Maxim	um possible scon	e using this molecu	le type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	1632	SEQ ID NO: 7921	0.09 %	3607.31448	
2	1640	SEQ ID NO: 7922	0.04 %	1748.2560912	
3	1776	SEQ ID NO: 7923	0.03 %	1492.58592	
4	2512	SEQ ID NO: 7924	0.03 %	1434.16845	
5	1073	SEQ ID NO: 7925	0.03 %	1338.876	
6	230	SEQ ID NO: 7926	0.01 %	685.78272	
7	1001	SEQ ID NO: 7927	0.01 %	559.8936	
8	716	SEQ ID NO: 7928	0.01 %	558.27486	

9	2280	SEQ ID NO: 7929	0.01 %	511.19781048
10	590	SEQ ID NO: 7930	0.01 %	469.6692
11	664	SEQ ID NO: 7931	0.01 %	442.076389524
12	1094	SEQ ID NO: 7932	0.00 %	382.536
13	1735	SEQ ID NO: 7933	0.00 %	382.536
14	1625	SEQ ID NO: 7934	0.00 %	342.4606344
15	1974	SEQ ID NO: 7935	0.00 %	336.885048
16	2382	SEQ ID NO: 7936	0.00 %	319.9392
_17	2417	SEQ ID NO: 7937	0.00 %	319.9392
18	744	SEQ ID NO: 7938	0.00 %	256.416670125
19	108	SEQ ID NO: 7939	0.00 %	232.52724
20	390	SEQ ID NO: 7940	0.00 %	228.0411084

HLA A 0201 - 10 mers				
Maxim	um possible sco	re using this mole	cule type	3925227.1
Rank	Start position	Sequence	% of max. score	Score
11	2511	SEQ ID NO: 7941	0.38 %	15126.90795
2	1608	SEQ ID NO: 7942	0.05 %	2049.4656
3	2572 '	SEQ ID NO: 7943	0.04 %	1879.5921264
4	255	SEQ ID NO: 7944	0.03 %	1566.6522795
5	895	SEQ ID NO: 7945	0.03 %	1338.876
6	1171	SEQ ID NO: 7946	0.02 %	1107.960876
7	1691	SEQ ID NO: 7947	0.01 %	782.95521024
8	20	SEQ ID NO: 7948	0.01 %	549.9372312
9	1632	SEQ ID NO: 7949	0.01 %	479.041993296
10	2280	SEQ ID NO: 7950	0.01 %	472.418344576987
11	1963	SEQ ID NO: 7951	0.00 %	358.73928
12	1955	SEQ ID NO: 7952	0.00 %	331.093464
13	741	SEQ ID NO: 7953	0.00 %	318.652488
14	523	SEQ ID NO: 7954	0.00 %	278.7876
15	1073	SEQ ID NO: 7955	0.00 %	266.6988828
16	2489	SEQ ID NO: 7956	0.00 %	243.432
17	777	SEQ ID NO: 7957	0.00 %	218.5730664
18	1737	SEQ ID NO: 7958	0.00 %	218.0785572
19	589	SEQ ID NO: 7959	0.00 %	210.538251
20	229	SEQ ID NO: 7960	0.00 %	205.230564

HLA A 1101 - 9 mers	
Maximum possible score using this molecule type	36

Rank	Start position	Sequence	% of max. score	Score
1	2337	SEQ ID NO: 7961	33.33 %	12
2	2156	SEQ ID NO: 7962	25 %	. 9
3	492	SEQ ID NO: 7963	20 %	7.2
4	18	SEQ ID NO: 7964	16.66 %	6
5	332	SEQ ID NO: 7965	16.66 %	6
6	415	SEQ ID NO: 7966	16.66 %	6
7	2479	SEQ ID NO: 7967	16.66 %	- 6
8	1495	SEQ ID NO: 7968	11.11 %	4
9	2035	SEQ ID NO: 7969	11.11 %	4
10	1349	SEQ ID NO: 7970	10 %	3.6
11	1194	SEQ ID NO: 7971	8.33 %	3
12	1648	SEQ ID NO: 7972	8.33 %	3
13	96	SEQ ID NO: 7973	6.66 %	2.4
14	764	SEQ ID NO: 7974	6.66 %	2.4
15	986	SEQ ID NO: 7975	6.66 %	2.4
16	2345	SEQ ID NO: 7976	6.66 %	2.4
17	698	SEQ ID NO: 7977	5.55 %	2
18	1355	SEQ ID NO: 7978	5.55 %	2
19	1987	SEQ ID NO: 7979	5.55 %	2
20	2085	SEQ ID NO: 7980	5.55 %	2

	HLA A 1101 - 10 mers				
Maximu	m possible score us	sing this molecule type		36	
Rank	Start position	Sequence	% of max. score	Score	
1	2083	SEQ ID NO: 7981	33.33 %	12	
2	2123	SEQ ID NO: 7982	25 %	9	
3	2147	SEQ ID NO: 7983	16.66 %	6	
4	331	SEQ ID NO: 7984	12.5 %	4.5	
5	1035	SEQ ID NO: 7985	11.11 %	4	
6	1064	SEQ ID NO: 7986	11.11 %	4	
7	2154	SEQ ID NO: 7987	11.11 %	4	
8	1048	SEQ ID NO: 7988	7.5 %	2.7	
9	202	SEQ ID NO: 7989	6.66 %	2.4	
10	721	SEQ ID NO: 7990	6.66 %	2.4	
11	2109	SEQ ID NO: 7991	6.66 %	2.4	
12	2230	SEQ ID NO: 7992	6.66 %	2.4	
13	1306	SEQ ID NO: 7993	5.55 %	2	
14	1622	SEQ ID NO: 7994	5.55 %	2	

15	1772	SEQ ID NO: 7995	5.55 %	2
16	1796	SEQ ID NO: 7996	5.55 %	2
17	186	SEQ ID NO: 7997	5 %	1.8
18	414	SEQ ID NO: 7998	5 %	1.8
19	697	SEQ ID NO: 7999	5 %	1.8
20	1175	SEQ ID NO: 8000	5 %	1.8

HLA B7 - 9 mers				
Maximu	m possible score us	sing this molecule type		5400
Rank	Start position	Sequence	% of max. score	Score
1	1447	SEQ ID NO: 8001	14.81 %	800
2	642	SEQ ID NO: 8002	3.70 %	200
3	34	SEQ ID NO: 8003	2.22 %	120
4	186	SEQ ID NO: 8004	1.48 %	1 80
5	244	SEQ ID NO: 8005	1.48 %	80
6	459	SEQ ID NO: 8006	1.48 %	80
7	1475	SEQ ID NO: 8007	1.48 %	. 80
8	1867	SEQ ID NO: 8008	1.48 %	80
9	2032	SEQ ID NO: 8009	1.48 %	80
10	2047	SEQ ID NO: 8010	1.48 %	80
11	2335	SEQ ID NO: 8011	1.48 %	80
12	622	SEQ ID NO: 8012	1.11 %	60
13	1375	SEQ ID NO: 8013	1.11 %	. 60
14	1617	SEQ ID NO: 8014	0.92 %	50
15	1023	SEQ ID NO: 8015	0.83 %	45
16	286	SEQ ID NO: 8016	0.74 %	. 40
17	490	SEQ ID NO: 8017	0.74 %	40
18	810	SEQ ID NO: 8018	0.74 %	40
19	1420	SEQ ID NO: 8019	0.74 %	40
20	1854	SEQ ID NO: 8020	0.74 %	40

	HLA B7 - 10 mers				
Maximu	m possible score us	sing this molecule type		5400	
Rank	Rank Start position Sequence % of max. score				
1	1617	SEQ ID NO: 8021	3.70 %	200	
2	752	SEQ ID NO: 8022	2.22 %	120	
3	1552	SEQ ID NO: 8023	2.22 %	120	
4	154	SEQ ID NO: 8024	1.48 %	80	
5	165	SEQ ID NO: 8025	1.48 %	80	

6	383	SEQ ID NO: 8026	1.48 %	80
7	1501	SEQ ID NO: 8027	1.48 %	80
8	2093	SEQ ID NO: 8028	1.48 %	80
9	2564	SEQ ID NO: 8029	1.48 %	80
10	622	SEQ ID NO: 8030	1.11 %	60
11	1086	SEQ ID NO: 8031	1.11 %	60
12	1262	SEQ ID NO: 8032	1.11 %	60
13	1556	SEQ ID NO: 8033	1.11 %	- 60 ·
14	845	SEQ ID NO: 8034	1 %	54
15	286	SEQ ID NO: 8035	0.74 %	40
16	490	SEQ ID NO: 8036	0.74 %	40
17	552	SEQ ID NO: 8037	0.74 %	40
18	1858	SEQ ID NO: 8038	0.74 %	40
19	2107	SEQ ID NO: 8039	0.74 %	40
20	2582	SEQ ID NO: 8040	0.74 %	40

Table 16: Epitopes for SEQ ID NO: 6042

		HLA A1 - 9 mers		
Maximu	m possible score us	sing this molecule type		5625
Rank	Start position	Sequence	% of max. score	Score
1	846	SEQ ID NO: 8041	2.22 %	125
2	798	SEQ ID NO: 8042	1.6 %	90
3	787	SEQ ID NO: 8043	0.88 %	50
4	1178	SEQ ID NO: 8044	0.88 %	50
5	637	SEQ ID NO: 8045	0.8 %	45
6	557	SEQ ID NO: 8046	0.44 %	25
7	1020	SEQ ID NO: 8047	0.44 %	25
8	282	SEQ ID NO: 8048	0.32 %	18
9	1241	SEQ ID NO: 8049	0.24 %	13.5
10	466	SEQ ID NO: 8050	0.22 %	12.5
11	727	SEQ ID NO: 8051	0.2 %	11.25
12	706	SEQ ID NO: 8052	0.17 %	10
13	324	SEQ ID NO: 8053	0.16 %	9
14	752	SEQ ID NO: 8054	0.16 %	9
15	54	SEQ ID NO: 8055	0.13 %	7.5
16	554	SEQ ID NO: 8056	0.13 %	7.5
17	590	SEQ ID NO: 8057	0.12 %	6.75

18	569	SEQ ID NO: 8058	0.08 %	5
19	613	SEQ ID NO: 8059	0.08 %	5
20	90	SEQ ID NO: 8060	0.08 %	4.5

	HLA A1 - 10 mers				
		sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1	1241	SEQ ID NO: 8061	4.8 %	-270	
2	967	SEQ ID NO: 8062	0.8 %	45	
3	1010	SEQ ID NO: 8063	0.48 %	27	
4	426	SEQ ID NO: 8064	0.44 %	25	
5	· 809	SEQ ID NO: 8065	0.44 %	25	
6	1178	SEQ ID NO: 8066	0.44 %	25	
7	787	SEQ ID NO: 8067	0.22 %	12.5	
8	958	SEQ ID NO: 8068	0.22 %	12.5	
. 9	727	SEQ ID NO: 8069	0.2 %	11.25	
10	610	SEQ ID NO: 8070	0.17 %	10	
11	12	SEQ ID NO: 8071	0.13 %	7.5	
12	1181	SEQ ID NO: 8072	0.12 %	6.75	
13	373	SEQ ID NO: 8073	0.11 %	6.25	
14	602	SEQ ID NO: 8074	0.11 %	6.25	
15	20	SEQ ID NO: 8075	0.04 %	2.5	
16	32	SEQ ID NO: 8076	0.04 %	2.5	
17	53	SEQ ID NO: 8077	0.04 %	2.5	
18	· 400	SEQ ID NO: 8078	0.04 %	2.5	
19	557	SEQ ID NO: 8079	0.04 %	2.5	
20	667	SEQ ID NO: 8080	0.04 %	2.5	

	HLA A3 - 9 mers				
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
11	768	SEQ ID NO: 8081	0.82 %	100	
2	808	SEQ ID NO: 8082	0.49 %	60	
3	85	SEQ ID NO: 8083	0.24 %	30	
4	663	SEQ ID NO: 8084	0.24 %	30	
5	1245	SEQ ID NO: 8085	0.14 %	18	
6	288	SEQ ID NO: 8086	0.09 %	12	
7	50	SEQ ID NO: 8087	0.08 %	10	
8	320	SEQ ID NO: 8088	0.07 %	9	

9	402	SEQ ID NO: 8089	0.07 %	9
10	798	SEQ ID NO: 8090	0.07 %	9
11	902	SEQ ID NO: 8091	0.06 %	8.1
12	364	SEQ ID NO: 8092	0.05 %	6:75
13	297	SEQ ID NO: 8093	0.04 %	6
14	992	SEQ ID NO: 8094	0.04 %	6
15	38	SEQ ID NO: 8095	0.03 %	4.5
16	249	SEQ ID NO: 8096	0.03 %	-4.5-
17	706	SEQ ID NO: 8097	0.03 %	4.05
18	1204	SEQ ID NO: 8098	0.03 %	4.05
19	1178	SEQ ID NO: 8099	0.03 %	4
20	343	SEQ ID NO: 8100	0.02 %	3.6

	HLA A3 - 10 mers			
Maximu	m possible score us	sing this molecule type		12150
Rank	Start position	Sequence	% of max. score	Score
. 1	255	SEQ ID NO: 8101	1.48 %	180
2	180	SEQ ID NO: 8102	0.55 %	67.5
3	768	SEQ ID NO: 8103	0.49 %	60
4	1177	SEQ ID NO: 8104	0.49 %	60
5	380	SEQ ID NO: 8105	0.24 %	30
6	100	SEQ ID NO: 8106	0.18 %	22.5
7	786	SEQ ID NO: 8107	0.16 %	20
8	1217	SEQ ID NO: 8108	0.16 %	20
9	207	SEQ ID NO: 8109	0.14 %	18
10	1183	SEQ ID NO: 8110	0.14 %	18
11	38	SEQ ID NO: 8111	0.09 %	12
12	52	SEQ ID NO: 8112	0.09 %	12
13	8	SEQ ID NO: 8113	0.06 %	8
14	679	SEQ ID NO: 8114	0.06 %	8
15	73	SEQ ID NO: 8115	0.05 %	6.75
16	1204	SEQ ID NO: 8116	0.05 %	6.075
17	50	SEQ ID NO: 8117	0.04 %	6
18	774	SEQ ID NO: 8118	0.04 %	6
19	845	SEQ ID NO: 8119	0.04 %	6
20	214	SEQ ID NO: 8120	0.04 %	5.4

HLA A24 - 9 mers	
Maximum possible score using this molecule type	1596.672

Rank	Start position	Sequence	% of max. score	Score
1	1118	SEQ ID NO: 8121	19.84 %	316.8
2	51	SEQ ID NO: 8122	18.78 %	300
3	161	SEQ ID NO: 8123	18.78 %	300
4	434	SEQ ID NO: 8124	18.78 %	300
5	365	SEQ ID NO: 8125	13.77 %	220
6	736	SEQ ID NO: 8126	12.52 %	200
- 7	620	SEQ ID NO: 8127	7.51 %	120
8	1068	SEQ ID NO: 8128	7.51 %	120
9	817	SEQ ID NO: 8129	3.75 %	60
10	336	SEQ ID NO: 8130	3.44 %	55
11	687	SEQ ID NO: 8131	3.13 %	50
12	254	SEQ ID NO: 8132	2.34 %	37.5
13	627	SEQ ID NO: 8133	1.87 %	30
14	950	SEQ ID NO: 8134	1.75 %	28
15	28	SEQ ID NO: 8135	1.56 %	25
16	408	SEQ ID NO: 8136	1.56 %	25
17	159	SEQ ID NO: 8137	1.31 %	21
18	1166	SEQ ID NO: 8138	1.26 %	20.16
19	45	SEQ ID NO: 8139	1.25 %	20
20	185	SEQ ID NO: 8140	1.25 %	20

HLA A24 - 10 mers				
Maximum possible score using this molecule type			ре	1596.672
Rank	Start position	Sequence	% of max. score	Score
:1	438	SEQ ID NO: 8141	27.55 %	440
2	489	SEQ ID NO: 8142	22.54 %	360
3	254	SEQ ID NO: 8143	18.78 %	300
4	354	SEQ ID NO: 8144	11.27 %	180
5	406	SEQ ID NO: 8145	11.27 %	180
6	1047	SEQ ID NO: 8146	11.27 %	180
7	473	SEQ ID NO: 8147	7.51 %	120
8	350	SEQ ID NO: 8148	6.26 %	100
9	769	SEQ ID NO: 8149	6.26 %	100
10	193	SEQ ID NO: 8150	5.63 %	90
11	479	SEQ ID NO: 8151	3.13 %	50
12	0	SEQ ID NO: 8152	2.70 %	43.2
13	813	SEQ ID NO: 8153	1.87 %	30
14	739	SEQ ID NO: 8154	1.50 %	24
		-421-		

15	782	SEQ ID NO: 8155	1.50 %	24
16	1186	SEQ ID NO: 8156	1.31 %	21
17	910	SEQ ID NO: 8157	1.05 %	16.8
18	128	SEQ ID NO: 8158	0.93 %	15
19	183	SEQ ID NO: 8159	0.93 %	15
20	1069	SEQ ID NO: 8160	0.93 %	15

	HLA A 0201 - 9 mers				
Maxim	um possible score	using this molecule	e type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	1041	SEQ ID NO: 8161	0.01 %	484.2379773	
2	981	SEQ ID NO: 8162	0.00 %	382,536	
3	957	SEQ ID NO: 8163	0.00 %	342.4606344	
4	896	SEQ ID NO: 8164	0.00 %	232.6931712	
5	1173	SEQ ID NO: 8165	0.00 %	201.447432	
6	733	SEQ ID NO: 8166	0.00 %	171.86796	
7	410	SEQ ID NO: 8167	0.00 %	135.45252	
8	786	SEQ ID NO: 8168	0.00 %	119.463012	
9	150	SEQ ID NO: 8169	0.00 %	102.17550222	
10	1	SEQ ID NO: 8170	0.00 %	94.98737754	
11	595	SEQ ID NO: 8171	0.00 %	93.239424	
12	1095	SEQ ID NO: 8172	0.00 %	89.41779	
13	1166	SEQ ID NO: 8173	0.00 %	87.58584	
14	845	SEQ ID NO: 8174	0.00 %	79.642008	
15	734	SEQ ID NO: 8175	0.00 %	73.47672	
16	802	SEQ ID NO: 8176	0.00 %	71.872056	
17.	1213	SEQ ID NO: 8177	0.00 %	71.872056	
18	105	SEQ ID NO: 8178	0.00 %	50.232	
19	939	SEQ ID NO: 8179	0.00 %	49.13352	
20	130	SEQ ID NO: 8180	0.00 %	48.732354	

	HLA A 0201 - 10 mers				
Maxim	um possible scor	e using this molecul	e type	3925227.1	
Rank	nk Start position Sequence % of max. score		Score		
1	372	SEQ ID NO: 8181	0.04 %	1896.33528	
2	410	SEQ ID NO: 8182	0.02 %	1134.00849744	
3	162	SEQ ID NO: 8183	0.01 %	685.3897512	
4	1076	SEQ ID NO: 8184	0.01 %	640.90320525	
5	1196	SEQ ID NO: 8185	0.01 %	623.742666372	

6	353	SEQ ID NO: 8186	0.01 %	446.7384576
7	50	SEQ ID NO: 8187	0.00 %	375.97824
8	733	SEQ ID NO: 8188	0.00 %	271.863864
9	130	SEQ ID NO: 8189	0.00 %	235.6873848
10	415	SEQ ID NO: 8190	0.00 %	185.679
11	297	SEQ ID NO: 8191	0.00 %	177.496704
12 .	1	SEQ ID NO: 8192	0.00 %	152.42160582
13	56	SEQ ID NO: 8193	0.00 %	110.013876
14	732	SEQ ID NO: 8194	0.00 %	101.0988
15	6	SEQ ID NO: 8195	0.00 %	98.26704
16	261	SEQ ID NO: 8196	0.00 %	91.60164
17	1040	SEQ ID NO: 8197	0.00 %	76.98537
18	928	SEQ ID NO: 8198	0.00 %	71.2908
19	1188	SEQ ID NO: 8199	0.00 %	69.81282
20	1094	SEQ ID NO: 8200	0.00 %	52.5987

	HLA A 1101 - 9 mers			
Maximu	m possible score us	sing this molecule type		36
Rank	Start position	Sequence	% of max. score	Score
11	402	SEQ ID NO: 8201	25 %	· 9
2	902	SEQ ID NO: 8202	22.5 %	8.1
· 3	288	SEQ ID NO: 8203	11.11 %	4
4	85	SEQ ID NO: 8204	6.66 %	2.4
5	706	SEQ ID NO: 8205	6.66 %	2.4
-6	456	SEQ ID NO: 8206	5.55 %	2
7	920	SEQ ID NO: 8207	5.55 %	2
8	535	SEQ ID NO: 8208	5 %	1.8
9	364	SEQ ID NO: 8209	3.33 %	1.2
10	438	SEQ ID NO: 8210	3.33 %	1.2
11	798	SEQ ID NO: 8211	3.33 %	1.2
12	808	SEQ ID NO: 8212	3.33 %	1.2
13	937	SEQ ID NO: 8213	[,] 3.33 %	1.2
14	956	SEQ ID NO: 8214	3.33 %	1.2
15	557	SEQ ID NO: 8215	2.77 %	1
16	1218	SEQ ID NO: 8216	2.77 %	1
17	784	SEQ ID NO: 8217	2.5 %	0.9
18	249	SEQ ID NO: 8218	2.22 %	0.8
_19	768	SEQ ID NO: 8219	2.22 %	0.8
20	1178	SEQ ID NO: 8220	2.22 %	0.8

	HLA A 1101 - 10 mers			
Maximu	m possible score us	sing this molecule type		36
Rank	Start position	Sequence	% of max. score	Score
1	38	SEQ ID NO: 8221	13.33 %	4.8
2	807	SEQ ID NO: 8222	12.5 %	4.5
3	100	SEQ ID NO: 8223	11.11 %	4
4	380	SEQ ID NO: 8224	11.11 %	- 4
5	767	SEQ ID NO: 8225	10 %	3.6
6	533	SEQ ID NO: 8226	8.33 %	3
7	967	SEQ ID NO: 8227	6.66 %	2.4
8	919	SEQ ID NO: 8228	5.55 %	2
9	305	SEQ ID NO: 8229	5 %	1.8
10	. 211	SEQ ID NO: 8230	3.33 %	1.2
11	511	SEQ ID NO: 8231	3.33 %	1.2
12	1177	SEQ ID NO: 8232	3.33 %	1.2
13	429	SEQ ID NO: 8233	2.77 %	1
14	758	SEQ ID NO: 8234	2.77 %	1
15	797	SEQ ID NO: 8235	2.5 %	0.9
16	255	SEQ ID NO: 8236	2.22 %	0.8
17	986	SEQ ID NO: 8237	2.22 %	0.8
18	1157	SEQ ID NO: 8238	2.22 %	. 0.8
19	170	SEQ ID NO: 8239	1.66 % :	0.6
20	893	SEQ ID NO: 8240	1.66 %	0.6

	HLA B7 - 9 mers			
Maximu	m possible score us	sing this molecule type	у	5400
Rank	Start position	Sequence	% of max. score	Score
1	200	SEQ ID NO: 8241	1.48 %	80
2	1243	SEQ ID NO: 8242	1.48 %	80
3	123	SEQ ID NO: 8243	0.74 %	40
4	248	SEQ ID NO: 8244	0.66 %	36
5	1036	SEQ ID NO: 8245	0.66 %	36
6	494	SEQ ID NO: 8246	0.37 %	20
7	495	SEQ ID NO: 8247	0.37 %	20
8	523	SEQ ID NO: 8248	0.37 %	20
9	842	SEQ ID NO: 8249	0.37 %	20
10	932	SEQ ID NO: 8250	0.37 %	20
11	274	SEQ ID NO: 8251	0.33 %	18

12	588	SEQ ID NO: 8252	0.22 %	12
13	656	SEQ ID NO: 8253	0.22 %	12
14	657	SEQ ID NO: 8254	0.22 %	12
15	767	SEQ ID NO: 8255	0.22 %	12
16	911	SEQ ID NO: 8256	0.22 %	12
17	939	SEQ ID NO: 8257	0.22 %	12
18	1007	SEQ ID NO: 8258	0.22 %	12
19	1170	SEQ ID NO: 8259	0.22 %	- 12
20	1206	SEQ ID NO: 8260	0.22 %	12

HLA B7 - 10 mers				
Maximu	m possible score us	sing this molecule type		5400
Rank	Start position	Sequence	% of max. score	Score
1	505	SEQ ID NO: 8261	4.44 %	240
2	312	SEQ ID NO: 8262	3.70 %	200
3	141	SEQ ID NO: 8263	1.11 %	60
4	1006	SEQ ID NO: 8264	0.66 %	. 36
[.] 5	411	SEQ ID NO: 8265	0.44 %	24
6	122	SEQ ID NO: 8266	0.37 %	20
7	134	SEQ ID NO: 8267	0.37 %	20
8	184	SEQ ID NO: 8268	0.37 %	20
9	367	SEQ ID NO: 8269	0.37 %	20
10	402	SEQ ID NO: 8270	0.37 %	20
11	494	SEQ ID NO: 8271	0.37 %	20
12	560	SEQ ID NO: 8272	0.37 %	20
13	626	SEQ ID NO: 8273	0.37 %	20
14	931	SEQ ID NO: 8274	0.37 %	20
15	956	SEQ ID NO: 8275	0.37 %	20
16	1117	SEQ ID NO: 8276	0.37 %	20
17	1169	SEQ ID NO: 8277	0.37 %	20
18	1196	SEQ ID NO: 8278	0.37 %	20
19	247	SEQ ID NO: 8279	0.22 %	12
20	273	SEQ ID NO: 8280	0.22 %	12

Table 17: Epitopes for SEQ ID NO: 6043

HLA A1 - 9 mers	
Maximum possible score using this molecule type	5625

Rank	Start position	Sequence	% of max. score	Score
1	168	SEQ ID NO: 8281	0.2 %	11.25
2	212	SEQ ID NO: 8282	0.08 %	4.5
3	223	SEQ ID NO: 8283	0.08 %	4.5
4	104	SEQ ID NO: 8284	0.04 %	2.5
5	170	SEQ ID NO: 8285	0.04 %	2.5
6	99	SEQ ID NO: 8286	0.04 %	2.25
7	188	SEQ ID NO: 8287	0.02 %	1.35
8	180	SEQ ID NO: 8288	0.02 %	1.25
9	219	SEQ ID NO: 8289	0.02 %	1.25
10	18	SEQ ID NO: 8290	0.01 %	1
11	226	SEQ ID NO: 8291	0.01 %	1
12	98	SEQ ID NO: 8292	0.01 %	0.625
13	151	SEQ ID NO: 8293	0.01 %	0.625
14	10	SEQ ID NO: 8294	0.01 %	0.6
15	13	SEQ ID NO: 8295	0.00 %	0.5
16	32	SEQ ID NO: 8296	0.00 %	0.5
17	70	SEQ ID NO: 8297	0.00 %	0.5
18	78	SEQ ID NO: 8298	0.00 %	0.5
19	82	SEQ ID NO: 8299	0.00 %	0.5
20	145	SEQ ID NO: 8300	0.00 %	0.5

	HLA A1 - 10 mers			
Maximu	m possible score us	ing this molecule type		5625
Rank	Start position	Sequence	% of max. score	Score
1	99	SEQ ID NO: 8301	0.8 %	45
2	223	SEQ ID NO: 8302	0.8 %	45
3	188	SEQ ID NO: 8303	0.48 %	27
4	206	SEQ ID NO: 8304	0.2 %	11.25
5	253	SEQ ID NO: 8305	0.17 %	10
6	174	SEQ ID NO: 8306	0.13 %	7.5
7	97	SEQ ID NO: 8307	0.04 %	2.5
8	257	SEQ ID NO: 8308	0.04 %	2.5
9	179	SEQ ID NO: 8309	0.04 %	2.25
10	162	SEQ ID NO: 8310	0.02 %	1.25
11	196	SEQ ID NO: 8311	0.02 %	1.25
12	219	SEQ ID NO: 8312	0.02 %	1.25
13	18	SEQ ID NO: 8313	0.01 %	1
14	246	SEQ ID NO: 8314	0.01 %	11

15	38	SEQ ID NO: 8315	0.01 %	0.75
16	33	SEQ ID NO: 8316	0.00 %	0.5
17	69	SEQ ID NO: 8317	0.00 %	0.5
18	81	SEQ ID NO: 8318	0.00 %	0.5
19	104	SEQ ID NO: 8319	0.00 %	0.5
20	116	SEQ ID NO: 8320	0.00 %	0.5

HLA A3 - 9 mers				
		sing this molecule type		12150
Rank	Start position	Sequence	% of max. score	Score
1	104	SEQ ID NO: 8321	0.98 %	120
2	123	SEQ ID NO: 8322	0.74 %	90
3	82	SEQ ID NO: 8323	0.44 %	54
4	106	SEQ ID NO: 8324	0.11 %	13.5
5	99	SEQ ID NO: 8325	0.08 %	10.8
6	127	SEQ ID NO: 8326	0.08 %	10
7	71	SEQ ID NO: 8327	0.07 %	9
8	1	SEQ ID NO: 8328	0.06 %	8.1
9	113	' SEQ ID NO: 8329	0.04 %	6
10	84	SEQ ID NO: 8330	0.03 %	4.5
11	109	SEQ ID NO: 8331	0.03 %	4.05
12	58	SEQ ID NO: 8332	0.02 %	3
13	138	SEQ ID NO: 8333	0.02 %	3
14	44	SEQ ID NO: 8334	0.02 %	2.7
_ 15	81	SEQ ID NO: 8335	0.02 %	2.7
16	226	SEQ ID NO: 8336	0.02 %	2.7
17	184	SEQ ID NO: 8337	0.01 %	1.8
18	102	SEQ ID NO: 8338	0.01 %	1.215
19	39	SEQ ID NO: 8339	0.00 %	1.2
20	234	SEQ ID NO: 8340	0.00 %	0.9

	HLA A3 - 10 mers				
Maximu	m possible score us	sing this molecule type		12150	
Rank	Start position	Sequence	% of max. score	Score	
_ 1	99	SEQ ID NO: 8341	1.33 %	162	
2	81	SEQ ID NO: 8342	0.44 %	54	
3	104	SEQ ID NO: 8343	0.24 %	30	
4	51	SEQ ID NO: 8344	0.16 %	20	
5	122	SEQ ID NO: 8345	0.11 %	13.5	

6	71	SEQ ID NO: 8346	0.07 %	9
7	69	SEQ ID NO: 8347	0.04 %	6
8	223	SEQ ID NO: 8348	0.04 %	5.4
9	84	SEQ ID NO: 8349	0.03 %	4.5
10	63	SEQ ID NO: 8350	0.02 %	3.6
11	138	SEQ ID NO: 8351	0.02.%	3
12	201	SEQ ID NO: 8352	0.01 %	1.8
13	44	SEQ ID NO: 8353	0.01,%	1.35
14	83	SEQ ID NO: 8354	0.01'%	1.35
15	116	SEQ ID NO: 8355	0.00 %	1.2
16	46	SEQ ID NO: 8356	0.00 %	0.9
.17	183	SEQ ID NO: 8357	0.00 %	0.81
18	57	SEQ ID NO: 8358	0.00 %	0.6
19	93	SEQ ID NO: 8359	0.00 %	0.6
20	113	SEQ ID NO: 8360	0.00 %	0.6

HLA A24 - 9 mers				
Maximu	Maximum possible score using this molecule type			1596.672
Rank	Start position	Sequence	% of max. score	Score
1	198	SEQ ID NO: 8361	13.15 %	210
2	105	SEQ ID NO: 8362	9.39 %	150
3	210	SEQ ID NO: 8363	4.69 %	75
4	75	SEQ ID NO: 8364	3.15 %	50.4
5	85	SEQ ID NO: 8365	2.63 %	42
6	205	SEQ ID NO: 8366	2.10 %	33.6
7	77	SEQ ID NO: 8367	1.87 %	30
8	158	SEQ ID NO: 8368	0.65 %	10.5
9	103	SEQ ID NO: 8369	0.56 %	9
10	227	SEQ ID NO: 8370	0.55 %	8.8704
11	32	SEQ ID NO: 8371	0.54 %	8.64
12	74	SEQ ID NO: 8372	0.50 %	8
13	131	SEQ ID NO: 8373	0.50 %	- 8
14	54	SEQ ID NO: 8374	0.46 %	7.5
15	99	SEQ ID NO: 8375	0.45 %	7.2
16	44	SEQ ID NO: 8376	0.37 %	6
17	62	SEQ ID NO: 8377	0.37 %	6
18	87	SEQ ID NO: 8378	0.37 %	6
19	89	SEQ ID NO: 8379	0.37 %	6
20	154	SEQ ID NO: 8380	0.37 %	6

HLA A24 - 10 mers				
	ım possible score ı	using this molecule ty	pe	1596.672
Rank	Start position	Sequence	% of max. score	Score
1	105	SEQ ID NO: 8381	22.54 %	360
2	204	SEQ ID NO: 8382	17.53 %	280
3	209	SEQ ID NO: 8383	3.13 %	50
4	75	SEQ ID NO: 8384	1.87 %	30
5	85	SEQ ID NO: 8385	1.87 %	30
6	77	SEQ ID NO: 8386	1.12 %	18
7	74	SEQ ID NO: 8387	0.84 %	13.44
8	210	SEQ ID NO: 8388	0.56 %	9
9	226	SEQ ID NO: 8389	0.55 %	8.8704
10	98	SEQ ID NO: 8390	0.54 %	8.64
11	198	SEQ ID NO: 8391	0.46 %	7.5
12	67	SEQ ID NO: 8392	0.45 %	7.2
13	152	SEQ ID NO: 8393	0.43 %	7
14	43	SEQ ID NO: 8394	0.37 %	6
. 15	63	SEQ ID NO: 8395	0.37 %	6
16	• 72	SEQ ID NO: 8396	0.37 %	6
17	89	SEQ ID NO: 8397	0.37 %	6
18	101	SEQ ID NO: 8398	0.37 %	6.
19	107	SEQ ID NO: 8399	0.37 %	6
20	111	SEQ ID NO: 8400	0.37 %	6

	HLA A 0201 - 9 mers				
		using this molecule	e type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	138	SEQ ID NO: 8401	0.21 %	8532.082944	
2	106	SEQ ID NO: 8402	0.10 %	3977.8497792	
3	44	SEQ ID NO: 8403	0.03 %	1243.078056	
4	71	SEQ ID NO: 8404	0.00 %	348.872832	
5	234	SEQ ID NO: 8405	0.00 %	243.432	
6	51	SEQ ID NO: 8406	0.00 %	130.26096	
7	109	SEQ ID NO: 8407	0.00 %	91.182672	
8	81	SEQ ID NO: 8408	0.00 %	73.342584	
9	88	SEQ ID NO: 8409	0.00 %	70.386624	
_10	11	SEQ ID NO: 8410	0.00 %	65.32728732	
11	38	SEQ ID NO: 8411	0.00 %	47.876409	

12	76	SEQ ID NO: 8412	0.00 %	36.8637882	
13	46	SEQ ID NO: 8413	0.00 %	30.889782	
14	211	SEQ ID NO: 8414	0.00 %	21.616753941	
15	201	SEQ ID NO: 8415	0.00 %	19.657134	
16	102	SEQ ID NO: 8416	0.00 %	18.4318941	
17	199	SEQ ID NO: 8417	0.00 %	16.496865	
18	74	SEQ ID NO: 8418	0.00 %	15.783256167	
19	62	SEQ ID NO: 8419	0.00 %	13.9968225	
20	99	SEQ ID NO: 8420	0.00 %	10.31851392	

HLA A 0201 - 10 mers							
Maxim	um possible scor	3925227.1					
Rank	Start position	Sequence	% of max. score	Score			
1	78	SEQ ID NO: 8421	0.01 %	556.494246			
2	138	SEQ ID NO: 8422	0.01 %	395.245972224			
3	84	SEQ ID NO: 8423	0.00 %	201.554244			
4	71	SEQ ID NO: 8424	0.00 %	143.65707264			
5	44	SEQ ID NO: 8425	0.00 %	132.54624			
6	· 76	SEQ ID NO: 8426	0.00 %	84.78671286			
7	8	SEQ ID NO: 8427	0.00 %	69.552			
8	211	SEQ ID NO: 8428	0.00 %	52.7237901			
9	113	SEQ ID NO: 8429	0.00 %	47.99088			
10	61	SEQ ID NO: 8430	0.00 %	37.4509575			
11	93	SEQ ID NO: 8431	0.00 %	31.24872			
12	137	SEQ ID NO: 8432	0.00 %	31.1384304			
13	. 37	SEQ ID NO: 8433	0.00 %	27.531			
14	55	SEQ ID NO: 8434	0.00 %	22.9153278			
15	98	SEQ ID NO: 8435	0.00 %	22.1063618985			
16	108	SEQ ID NO: 8436	0.00 %	21.55457052			
17	63	SEQ ID NO: 8437	0.00 %	21.3624			
18	45	SEQ ID NO: 8438	0.00 %	19.657134			
19	200	SEQ ID NO: 8439	0.00 %	19.657134			
20	104	SEQ ID NO: 8440	0.00 %	13.87622016			

HLA A 1101 - 9 mers							
Maximum possible score using this molecule type							
Rank	Start position	Sequence	% of max. score	Score			
1	58	SEQ ID NO: 8441	5.55 %	2			
2	125	SEQ ID NO: 8442	1.66 %	0.6			

3	226	SEQ ID NO: 8443	1.66 %	0.6
4	229	SEQ ID NO: 8444	1.66 %	0.6

	HLA A 1101 - 10 mers				
Maximum possible score using this molecule type				36	
Rank	Start position	Sequence	% of max. score	Score	
11	122	SEQ ID NO: 8445	2.22 %	0.8	
2	228	SEQ ID NO: 8446	2.22 %	- 0.8	

	HLA B7 - 9 mers				
Maximu	m possible score us	sing this molecule type		5400	
Rank	Start position	Sequence	% of max. score	Score	
1	97	SEQ ID NO: 8447	0.66 %	. 36	
2	86	SEQ ID NO: 8448	0.37 %	20	
3	37	SEQ ID NO: 8449	0.33 %	18	
4 .	62	SEQ ID NO: 8450	0.33 %	18	
5	32	SEQ ID NO: 8451	0.22 %	12	
6	102	SEQ ID NO: 8452	0.22 %	: 12	
7	227	SEQ ID NO: 8453	0.22 %	12	
8	53	SEQ ID NO: 8454	0.11 %	6	
9	1 .	SEQ ID NO: 8455	0.07 %	4	
10	44	SEQ ID NO: 8456	0.07 %	4	
11	56	SEQ ID NO: 8457	0.07 %	4	
12	64	SEQ ID NO: 8458	0.07 %	4	
13	74	SEQ ID NO: 8459	0.07 %	4	
14	76	SEQ ID NO: 8460	0.07 %	4	
15	87	SEQ ID NO: 8461	0.07 %	4	
16	106	SEQ ID NO: 8462	0.07 %	4	
17	131 '	SEQ ID NO: 8463	0.07 %	4	
18	23	SEQ ID NO: 8464	0.03 %	2	
19	157	SEQ ID NO: 8465	0.03 %	2	
20	166	SEQ ID NO: 8466	0.03 %	2	

	HLA B7 - 10 mers				
Maximum possible score using this molecule type					
Rank	Start position	Sequence	% of max. score	Score	
1	119	SEQ ID NO: 8467	3.33 %	180	
2	264	SEQ ID NO: 8468	1.48 %	80	
3	98	SEQ ID NO: 8469	0.66 %	36	

4	27	SEQ ID NO: 8470	0.37 %	20
5	86	SEQ ID NO: 8471	0.37 %	20
6	31	SEQ ID NO: 8472	0.22 %	12
7	63	SEQ ID NO: 8473	0.22'%	12
8	96	SEQ ID NO: 8474	0.22 %	12
9	101	SEQ ID NO: 8475	0.22 %	12
10	226	SEQ ID NO: 8476	0.22 %	12
11	157	SEQ ID NO: 8477	0.14 %	- 8 -
12	176	SEQ ID NO: 8478	0.14 %	8
13	238	SEQ ID NO: 8479	0.14 %	8
14	36	SEQ ID NO: 8480	0.11 %	6
15	53	SEQ ID NO: 8481	0.11 %	6
16	61	SEQ ID NO: 8482	0.11 %	6
17	3	SEQ ID NO: 8483	0.07 %	4
18	40	SEQ ID NO: 8484	0.07 %	4
19	55	SEQ ID NO: 8485	0.07 %	• 4
20	74	SEQ ID NO: 8486	0.07 %	4

Table 18: Epitopes for SEQ ID NO: 6044

	HLA A1 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	69	SEQ ID NO: 8487	0.04 %	2.5	
2	89	SEQ ID NO: 8488	0.02 %	1.5	
3	141	SEQ ID NO: 8489	0.01 %	1	
4	113	SEQ ID NO: 8490	0.00 %	0.5	

	HLA A1 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
11	21	SEQ ID NO: 8491	0.02 %	1.5	
2	88	SEQ ID NO: 8492	0.02 %	1.5	
3	8	SEQ ID NO: 8493	0.02 %	1.25	
4	31	SEQ ID NO: 8494	0.00 %	0.5	
5	112	SEQ ID NO: 8495	0.00 %	0.5	

HLA A3 - 9 mers

Maximu	m possible score us	sing this molecule type		12150
Rank	Start position	Sequence	% of max. score	Score
11	60	SEQ ID NO: 8496	1.23 %	150
2	77	SEQ ID NO: 8497	1.11 %	135
3	141	SEQ ID NO: 8498	0.49 %	60
4	95	SEQ ID NO: 8499	0.32 %	40
5	128	SEQ ID NO: 8500	0.08 %	10
6	113	SEQ ID NO: 8501	0.04 %	- 6
_ 7	69	SEQ ID NO: 8502	0.01 %	2
8	22	SEQ ID NO: 8503	0.01 %	1.8
9	42	SEQ ID NO: 8504	0.01 %	1.8
10	78 ·	SEQ ID NO: 8505	0.00 %	1.2
11	32	SEQ ID NO: 8506	0.00 %	1
12	54	SEQ ID NO: 8507	0.00 %	0.9
13	74	SEQ ID NO: 8508	0.00 %	0.9
14	28	SEQ ID NO: 8509	0.00 %	0.6
15	36	SEQ ID NO: 8510	0.00 %	0.6
16	48	SEQ ID NO: 8511	0.00 %	0.6
17	118	SEQ ID NO: 8512	0.00 %	0.6
18	4	SEQ ID NO: 8513	0.00 %	0.5

HLA A3 - 10 mers					
	m possible score us	sing this molecule type		12150	
Rank	Start position	Sequence	% of max. score	Score	
11	94	SEQ ID NO: 8514	0.49 %	60	
_ 2	48	SEQ ID NO: 8515	0.16 %	20	
3	128	SEQ ID NO: 8516	0.16 %	20 ·	
4	60	SEQ ID NO: 8517	0.12 %	15	
5	127	SEQ ID NO: 8518	0.12 %	15	
6	25	SEQ ID NO: 8519	0.04 %	6	
7	95	SEQ ID NO: 8520	0.04 %	6	
8	141	SEQ ID NO: 8521	0.04 %	6	
9	41	SEQ ID NO: 8522	0.04 %	5.4	
10	77	SEQ ID NO: 8523	0.04 %	5.4	
11	116	SEQ ID NO: 8524	0.04 %	5.4	
12	91	SEQ ID NO: 8525	0.03 %	4	
13	4	SEQ ID NO: 8526	0.01 %	2	
14	112	SEQ ID NO: 8527	0.01 %	1.8	
15	113	SEQ ID NO: 8528	0.01 %	1.35	

16	12	SEQ ID NO: 8529	0.00 %	1.2
17	31	SEQ ID NO: 8530	0.00 %	_ 1
18	32	SEQ ID NO: 8531	0.00 %	1
19	15	SEQ ID NO: 8532	0.00 %	0.9
20	27	SEQ ID NO: 8533	0.00 %	0.9

	HLA A24 - 9 mers				
Maximu	ım possible score ı	using this molecule ty	pe	1596.672	
Rank	Start position	Sequence	% of max. score	Score	
1	61	SEQ ID NO: 8534	14.46 %	231	
2	16	SEQ ID NO: 8535	3.13 %	50	
3 .	120	SEQ ID NO: 8536	1.87 %	30	
4	41	SEQ ID NO: 8537	0.60 %	9.6	
5	71	SEQ ID NO: 8538	0.45 %	7.2	
6	21	SEQ ID NO: 8539	0.37 %	6	
7	. 53	SEQ ID NO: 8540	0.37 %	6	
8	65 -	SEQ ID NO: 8541	0.37 %	- 6	
9	121	SEQ ID NO: 8542	0.37 %	6	
10	74	SEQ ID NO: 8543	0.36 %	5.76	
11	20	SEQ ID NO: 8544	0.35 %	5.6	
12	. 79	SEQ ID NO: 8545	0.35 %	5.6	
13	105	SEQ ID NO: 8546	0.33 %	5.28	
14	48	SEQ ID NO: 8547	0.30 %	4.8	
15	88	SEQ ID NO: 8548	0.30 %	4.8	
16	106	SEQ ID NO: 8549	0.30 %	4.8	
17	37	SEQ ID NO: 8550	0.27 %	4.4	
18	70	SEQ ID NO: 8551	0.27 %	4.4	
19	18	SEQ ID NO: 8552	0.25 %	4	
20	57	SEQ ID NO: 8553	0.22 %	3.6	

	HLA A24 - 10 mers				
Maximu	ım possible score ı	using this molecule ty	rpe	1596.672	
Rank	Start position	Sequence	% of max. score	Score	
1	120	SEQ ID NO: 8554	1.87 %	30	
2	73	SEQ ID NO: 8555	0.54 %	8.64	
3	19	SEQ ID NO: 8556	0.52 %	8.4	
4	78	SEQ ID NO: 8557	0.52 %	8.4	
5	104	SEQ ID NO: 8558	0.49 %	7.92	
6	61	SEQ ID NO: 8559	0.46 %	7.5	

	· 47	SEQ ID NO: 8560	0.45 %	7.2
8	36	SEQ ID NO: 8561	0.41 %	6.6
9	52	SEQ ID NO: 8562	0.37 %	6
10	64	SEQ ID NO: 8563	0.30 %	4.8
11	70	SEQ ID NO: 8564	0.30 %	4.8
12	105	SEQ ID NO: 8565	0.30 %	4.8
13	123	SEQ ID NO: 8566	0.30 %	4.8
14	69	SEQ ID NO: 8567	0.27 %	4.4
15	20	SEQ ID NO: 8568	0.25 %	4
16	66	SEQ ID NO: 8569	0.25 %	4
17	83	SEQ ID NO: 8570	0.25 %	4
18	86	SEQ ID NO: 8571	0.25 %	4
19	101	SEQ ID NO: 8572	0.25 %	: 4
20	119	SEQ ID NO: 8573	0.25 %	¹ 4

HLA A 0201 - 9 mers				
Maxim	um possible score	using this molecule	e type	3925227.1
Rank	Start position	Sequence	% of max. score	Score
1	62	SEQ ID NO: 8574	0.00 %	136.1646
2	85	SEQ ID NO: 8575	0.00 %	69.6969
3	47	SEQ ID NO: 8576	0.00 %	60.153786
4	121	SEQ ID NO: 8577	0.00 %	52.5182736
- 5	74	SEQ ID NO: 8578	0.00 %	49.13352
6	23	SEQ ID NO: 8579	0.00 %	21.99582
7	78	SEQ ID NO: 8580	0.00 %	19.42488
8	114	SEQ ID NO: 8581	0.00 %	14.6900655
9	4	SEQ ID NO: 8582	0.00 %	11.304684
10	79	SEQ ID NO: 8583	0.00 %	8.4687081
11	122	SEQ ID NO: 8584	0.00 %	6.0996
12	100	SEQ ID NO: 8585	0.00 %	5.382
13	105	SEQ ID NO: 8586	0.00 %	4.981593
14	25	SEQ ID NO: 8587	0.00 %	4.968
15	115	SEQ ID NO: 8588	0.00 %	4.966482
16	24	SEQ ID NO: 8589	0.00 %	4.4815221585
17	111	SEQ ID NO: 8590	0.00 %	4.128201
18	94	SEQ ID NO: 8591	0.00 %	3.67632
19	34	SEQ ID NO: 8592	0.00 %	3.47553
20	12	SEQ ID NO: 8593	0.00 %	3.30993

	HLA A 0201 - 10 mers				
Maxim	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	77	SEQ ID NO: 8594	0.00 %	147.97188	
2	. 62	SEQ ID NO: 8595	0.00 %	143.59176	
3	113	SEQ ID NO: 8596	0.00 %	106.83684	
4	78	SEQ ID NO: 8597	0.00 %	83.526984	
-5	86	SEQ ID NO: 8598	0.00 %	83.526984	
6	74	SEQ ID NO: 8599	0.00 %	69.552	
7	121	SEQ ID NO: 8600	0.00 %	61.06776	
8	12	SEQ ID NO: 8601	0.00 %	50.232	
9	44	SEQ ID NO: 8602	0.00 %	26.082	
10	4	SEQ ID NO: 8603	0.00 %	18.3816	
11	0	SEQ ID NO: 8604	0.00 %	17.38386	
12	72	SEQ ID NO: 8605	0.00 %	17.1396	
13	22	SEQ ID NO: 8606	0.00 %	16.21914	
14	122	SEQ ID NO: 8607	0.00 %	14.02908	
15	64	SEQ ID NO: 8608	0.00 %	11.161854	
16	46	SEQ ID NO: 8609	0.00 %	10.34586	
17	54	SEQ ID NO: 8610	0.00 %	8.846145	
18	47	SEQ ID NO: 8611	0.00 %	7.575080337	
19	131	SEQ ID NO: 8612	0.00 %	7.452	
20	114	SEQ ID NO: 8613	0.00 %	6.735366	

	HLA A 1101 - 9 mers				
Maximu	m possible score us	sing this molecule type		36	
Rank	Start position	Sequence	% of max. score	Score	
1	69	SEQ ID NO: 8614	5.55 %	2	
2	22	SEQ ID NO: 8615	5 %	1.8	
3	77	SEQ ID NO: 8616	5 %	1.8	
4	141	SEQ ID NO: 8617	3.33 %	1.2	
5	60	SEQ ID NO: 8618	2.22 %	0.8	
6	95	SEQ ID NO: 8619	2.22 %	0.8	
. 7	36	SEQ ID NO: 8620	1.66 %	0.6	

HLA A 1101 - 10 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	41	SEQ ID NO: 8621	3.33 %	1.2	

2	68	SEQ ID NO: 8622	3.33 %	1.2
3	94	SEQ ID NO: 8623	3.33 %	1.2
4	31	SEQ ID NO: 8624	2.77 %	1
5	127	SEO ID NO: 8625	2.5 %	0.9

HLA B7 - 9 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	48	SEQ ID NO: 8626	0.74 %	40	
2	20	SEQ ID NO: 8627	0.37 %	20	
3	121	SEQ ID NO: 8628	0.33 %	18	
4	18	SEQ ID NO: 8629	0.07 %	4	
5	21	SEQ ID NO: 8630	0.07 % !	4	
6	37	SEQ ID NO: 8631	0.07 %	4	
7	41	SEQ ID NO: 8632	0.07 %	4	
8	-53	SEQ ID NO: 8633	0.07 %	4	
9	65	SEQ ID NO: 8634	0.07 %	4	
10	70	SEQ ID NO: 8635	0.07 %	4	
11	71	SEQ ID NO: 8636	0.07 %	4	
12	74	SEQ ID NO: 8637	0.07 %	4	
13	79	SEQ ID NO: 8638	0.07 %	4	
14	88	SEQ ID NO: 8639	0.07 %	4	
15	105	SEQ ID NO: 8640	0.07 %	4	
16	106	SEQ ID NO: 8641	0.07 %	4	
17	124	SEQ ID NO: 8642	0.07 %	4	
18	1 ,	SEQ ID NO: 8643	0.03 %	- 2	
19	120	SEQ ID NO: 8644	0.03 %	1.8	
20	11	SEQ ID NO: 8645	0.02 %	1.2	

	HLA B7 - 10 mers				
	m possible score us	sing this molecule type		5400	
Rank	Start position	Sequence	% of max. score	Score	
1	66	SEQ ID NO: 8646	1.48 %	80	
2	123	SEQ ID NO: 8647	0.74 %	40	
3	20	SEQ ID NO: 8648	0.37 %	20	
4	64	SEQ ID NO: 8649	0.22 %	12	
5	119	SEQ ID NO: 8650	0.11 %	6	
6	54	SEQ ID NO: 8651	0.09 %	5	
7	19	SEQ ID NO: 8652	0.07 %	4	

8	36	SEQ ID NO: 8653	0.07 %	4
9	47	SEQ ID NO: 8654	0.07 %	4
10	52	SEQ ID NO: 8655	0.07 %	4
11	69	SEQ ID NO: 8656	0.07 %	4
12	70	SEQ ID NO: 8657	0.07 %	4
13	73	SEQ ID NO: 8658	0.07.%	4
14	78	SEQ ID NO: 8659	0.07.%	4
15	83	SEQ ID NO: 8660	0.07 %	- 4
16	86	SEQ ID NO: 8661	0.07 %	4
17	101	SEQ ID NO: 8662	0.07 %	4
18	104	SEQ ID NO: 8663	0.07 %	4
19	105	SEQ ID NO: 8664	0.07 %	4
20	15	SEQ ID NO: 8665	0.03 %	2

Table 19: Epitopes for SEQ ID NO: 6045

	HLA A1 - 9 mers				
Maximu	m possible score us	sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1	4	SEQ ID NO: 8666	0.02 %	1.35	
2	. 66	SEQ ID NO: 8667	0.02 %	1.35	
-3	- 33	SEQ ID NO: 8668	0.02 %	1.25	
4	. 44	SEQ ID NO: 8669	0.01 %	1	
5	50	SEQ ID NO: 8670	0.01 %	1	
6	14	SEQ ID NO: 8671	0.01 %	0.75	
7	48	SEQ ID NO: 8672	0.01 %	0.75	
8	11	SEQ ID NO: 8673	0.00 %	0.5	

	HLA A1 - 10 mers				
Maximu	m possible score us	sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1	4	SEQ ID NO: 8674	0.12 %	6.75	
2	· 66	SEQ ID NO: 8675	0.12 %	6.75	
3	10	SEQ ID NO: 8676	0.00 %	0.5	
4	28	SEQ ID NO: 8677	0.00 %	0.5	
5	32	SEQ ID NO: 8678	0.00 %	0.5	
6	47	SEQ ID NO: 8679	0.00 %	0.5	

	HLA A3 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
11	17	SEQ ID NO: 8680	0.24 %	30	
2	44	SEQ ID NO: 8681	0.07 %	9	
3	19	SEQ ID NO: 8682	0.06 %	8.1	
4	50	SEQ ID NO: 8683	0.04 %	5.4	
5	. 29	SEQ ID NO: 8684	0.03 %	- 4	
6	52	SEQ ID NO: 8685	0.02 %	3.24	
7	54	SEQ ID NO: 8686	0.02 %	3	
8	11	SEQ ID NO: 8687	0.01 %	1.8	
9 .	37	SEQ ID NO: 8688	0.01 %	1.8	
10	25	SEQ ID NO: 8689	0.01 %	1.35	
11	10	SEQ ID NO: 8690	0.00 %	0.9	
12	16	SEQ ID NO: 8691	0.00 %	0.9	
13	35	SEQ ID NO: 8692	0.00 %	0.6	

	HLA A3 - 10 mers					
Maximu	Maximum possible score using this molecule type					
Rank	Start position	Sequence	12150 Score			
1	49	SEQ ID NO: 8693	0.44 %	54		
2	17	SEQ ID NO: 8694	0.22 %	27		
' 3	10	SEQ ID NO: 8695	0.14 %	18		
4	16	SEQ ID NO: 8696	0.07 %	9		
5	32 -	SEQ ID NO: 8697	0.04 %	6		
6	19	SEQ ID NO: 8698	0.01 %	1.8		
7	29	SEQ ID NO: 8699	0.00 %	1.2		
8	23	SEQ ID NO: 8700	0.00 %	0.9		
9	26	SEQ ID NO: 8701	0.00 %	0.9		

	HLA A24 - 9 mers					
Maximu	Maximum possible score using this molecule type					
Rank	Start position	Sequence	% of max. score	Score		
1_1_	18	SEQ ID NO: 8702	1.87 %	30		
2	24	SEQ ID NO: 8703	0.65 %	10.5		
3	9	SEQ ID NO: 8704	0.52 %	8.4		
4	12	SEQ ID NO: 8705	0.52 %	8.4		
5	28	SEQ ID NO: 8706	0.52 %	8.4		
6	42	SEQ ID NO: 8707	0.52 %	8.4		

7	57	SEQ ID NO: 8708	0.52 %	8.4
8	66	SEQ ID NO: 8709	0.52 %	8.4
9	55	SEQ ID NO: 8710	0.51 %	8.25
10	0	SEQ ID NO: 8711	0.48 %	7.7
11	22	SEQ ID NO: 8712	0.45 %	7.2
12	10	SEQ ID NO: 8713	0.37 %	6
13	25	SEQ ID NO: 8714	0.37 %	- 6
14	30	SEQ ID NO: 8715	0.37 %	6
15	19	SEQ ID NO: 8716	0.35 %	5.6
16	40	SEQ ID NO: 8717	0.31 %	5
17	3	SEQ ID NO: 8718	0.30 %	4.8
18	65	SEQ ID NO: 8719	0.30 %	4.8
19	14	SEQ ID NO: 8720	0.27 %	4.32
20	56	SEQ ID NO: 8721	0.25 %	4

	HLA A24 - 10 mers				
Maximu	ım possible score ı	ising this molecule ty	rpe ;	1596.672	
Rank	Start position	Sequence	% of max. score	Score	
11	55	SEQ ID NO: 8722	18.78 %	300	
2	18	SEQ ID NO: 8723	2.63 %	42	
3	21	SEQ ID NO: 8724	2.25 %	36	
4	2	SEQ ID NO: 8725	1.87 %	30	
5	24	SEQ ID NO: 8726	1.87 %	30	
6	11	SEQ ID NO: 8727	0.52 %	8.4	
7	40	SEQ ID NO: 8728	0.52 %	8.4	
8	65	SEQ ID NO: 8729	0.42 %	6.72	
9	9	SEQ ID NO: 8730	0.37 %	6	
10	8	SEQ ID NO: 8731	0.35 %	5.6	
11	27	SEQ ID NO: 8732	0.35 %	5.6	
12	41	SEQ ID NO: 8733	0.35 %	5.6	
13	57	SEQ ID NO: 8734	0.31 %	5	
14	17	SEQ ID NO: 8735	0.25 %	4	
15	29	SEQ ID NO: 8736	0.25 %	4	
16	64	SEQ ID NO: 8737	0.25 %	4	
17	16	SEQ ID NO: 8738	0.22 %	3.6	
18	10	SEQ ID NO: 8739	0.18 %	3	
19	13	SEQ ID NO: 8740	0.18 %	2.88	
20	23	SEQ ID NO: 8741	0.08 %	1.4	

	HLA A 0201 - 9 mers				
Maxim	um possible scor	e using this molecul	e type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	19	SEQ ID NO: 8742	0.03 %	1310.8823136	
2	15	SEQ ID NO: 8743	0.02 %	1082.4143022	
3	16	SEQ ID NO: 8744	0.02 %	1040.33238624	
4	49	SEQ ID NO: 8745	0.00 %	382.536	
5	25	SEQ ID NO: 8746	0.00 %	342.863529264	
6	56	SEQ ID NO: 8747	0.00 %	63.28397376	
7	12	SEQ ID NO: 8748	0.00 %	40.19736105	
8	10	SEQ ID NO: 8749	0.00 %	21.3624	
9	22	SEQ ID NO: 8750	0.00 %	19.7762418	
10	26	SEQ ID NO: 8751	0.00 %	12.6684	
11	20	SEQ ID NO: 8752	0.00 %	11.544666	
12	37	SEQ ID NO: 8753	0.00 %	10.4328	
13	32	SEQ ID NO: 8754	0.00 %	8.4456	
14	23	SEQ ID NO: 8755	0.00 %	6.2888049	
15	47	SEQ ID NO: 8756	0.00 %	6.0858	
16	3 ,	SEQ ID NO: 8757	0.00 %	4.582929078	
17	18	SEQ ID NO: 8758	0.00 %	4.4855150505	
18	28	SEQ ID NO: 8759	0.00 %	4.2923589	
19	62	SEQ ID NO: 8760	0.00 %	2.88098391	
20	27	SEQ ID NO: 8761	0.00 %	1.699677	

	HLA A 0201 - 10 mers				
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	17	SEQ ID NO: 8762	0.16 %	6459.14167272	
2	19	SEQ ID NO: 8763	0.01 %	607.88448	
3	25	SEQ ID NO: 8764	0.00 %	126.83304	
4	11	SEQ ID NO: 8765	0.00 %	63.16728165	
5	15	SEQ ID NO: 8766	0.00 %	53.54651988	
6	37	SEQ ID NO: 8767	0.00 %	28.51632	
7	14	SEQ ID NO: 8768	0.00 %	21.8247414	
8	29	SEQ ID NO: 8769	0.00 %	21.3624	
9	26	SEQ ID NO: 8770	0.00 %	19.42488	
10	3	SEQ ID NO: 8771	0.00 %	17.2167282	
11	48	SEQ ID NO: 8772	0.00 %	15.7068219	
12	12	SEQ ID NO: 8773	0.00 %	9.8581266	

13	27	SEQ ID NO: 8774	0.00 %	7.3086111
14	39	SEQ ID NO: 8775	0.00 %	7.10976
15	23	SEQ ID NO: 8776	0.00 %	5.7419523
16	22	SEQ ID NO: 8777	0.00 %	4.599126
17	45	SEQ ID NO: 8778	0.00 %	2.5495155
18	31	SEQ ID NO: 8779	0.00 %	2.52747
19	52	SEQ ID NO: 8780	0.00 %	2.383605
20	20	SEQ ID NO: 8781	0.00 %	2.332847151

	HLA A 1101 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	44	SEQ ID NO: 8782	3.33 %	1.2	

HLA A 1101 - 10 mers				
Maximum possible score using this molecule type			36	
Rank	Start position	Sequence	% of max. score	Score

HLA B7 - 9 mers				
Maximu	ım possible score us	sing this molecule type		5400
Rank	Start position	Sequence	% of max. score	Score
1	3	SEQ ID NO: 8783	0.37 %	-20
2	12	SEQ ID NO: 8784	0.37 %	20
3	. 22	SEQ ID NO: 8785	0.37 %	20
4	56	SEQ ID NO: 8786	0.37 %	20
5	30	SEQ ID NO: 8787	0.22.%	12
6	9	SEQ ID NO: 8788	0.07 %	4
7	10	SEQ ID NO: 8789	0.07 %	4
8	19	SEQ ID NO: 8790	0.07 %	4
9	25	SEQ ID NO: 8791	0.07 %	4
10	28	SEQ ID NO: 8792	0.07 %	4
11	42	SEQ ID NO: 8793	0.07 %	4
12	65	SEQ ID NO: 8794	0.07 %	4
13	35	SEQ ID NO: 8795	0.05 %	3
14	66	SEQ ID NO: 8796	0.02 %	1.2
15	15	SEQ ID NO: 8797	0.01 %	1
16	47	SEQ ID NO: 8798	0.01 %	1
17	20	SEQ ID NO: 8799	0.01 %	0.6
18	23	SEQ ID NO: 8800	0.00 %	0.5

1 10 1	27 050 70	NO. 0004	
119.11	2/ SEO ID	NO: 8801 0.00	10/6 11 0 5 11
			, , o . o . o . o

HLA B7 - 10 mers				
Maximu	m possible score us	sing this molecule type)	5400
Rank	Start position	Sequence	% of max. score	Score
11	27	SEQ ID NO: 8802	0.37 %	20
2	8	SEQ ID NO: 8803	0.07 %	4
3	9	SEQ ID NO: 8804	0.07 %	- 4
4	11	SEQ ID NO: 8805	0.07 %	4
5	17	SEQ ID NO: 8806	0.07 %	4
6	29	SEQ ID NO: 8807	0.07 %	4
7	41	SEQ ID NO: 8808	0.07 %	4
8	52	SEQ ID NO: 8809	0.07 %	4
9	64	SEQ ID NO: 8810	0.07 %	4
_10	65	SEQ ID NO: 8811	0.07 %	4
11	3	SEQ ID NO: 8812	0.03 %	. 2
12	23	SEQ ID NO: 8813	0.03 %	2
13	21	SEQ ID NO: 8814	0.02 %	1.2
14	15	SEQ ID NO: 8815	0.01 %	1
15	35	SEQ ID NO: 8816	0.01 %	0.6
16	39	SEQ ID NO: 8817	0.01 %	0.6
17	12	SEQ ID NO: 8818	0.00 %	0.5
18	22	SEQ ID NO: 8819	0.00 %	0.5
19	45	SEQ ID NO: 8820	0.00 %	0.5

Table 20: Epitopes for SEQ ID NO: 6046

HLA A1 - 9 mers					
Maximu	m possible score u	sing this molecule type		5625	
Rank					
1	186	SEQ ID NO: 8821	2.22 %	125	
2	156	SEQ ID NO: 8822	0.88 %	50	
3	14	SEQ ID NO: 8823	0.08 %	4.5	
4	0	SEQ ID NO: 8824	0.04 %	2.5	
5	29	SEQ ID NO: 8825	0.04 %	2.5	
6	85	SEQ ID NO: 8826	0.04 %	2.5	
7	168	SEQ ID NO: 8827	0.04 %	2.5	
8	133	SEQ ID NO: 8828	0.02 %	1.35	

9	111	SEQ ID NO: 8829	0.02 %	1.125
10	61	SEQ ID NO: 8830	0.01 %	1
11	7	SEQ ID NO: 8831	0.01 %	0.9
12	131	SEQ ID NO: 8832	0.01 %	0.9
13	211	SEQ ID NO: 8833	0.01 %	0.625
14	4	SEQ ID NO: 8834	0.00 %	0.5
15	43	SEQ ID NO: 8835	0.00 %	0.5
16	95	SEQ ID NO: 8836	0.00 %	-0.5
17	136	SEQ ID NO: 8837	0.00 %	0.5

	HLA A1 - 10 mers				
Maximu	m possible score us	sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1	133 ,	SEQ ID NO: 8838	0.04 %	2.7	
2	84	SEQ ID NO: 8839	0.04 %	2.5	
3	167	SEQ ID NO: 8840	0.04 %	2.5	
4	186	SEQ ID NO: 8841	0.04 %	2.5	
5	131	SEQ ID NO: 8842	0.04 %	2.25	
6	14	SEQ ID NO: 8843	0.03 %	1.8	
7	205	SEQ ID NO: 8844	0.02 %	1.25	
8	111	SEQ ID NO: 8845	0.02 %	1.125	
9	60	SEQ ID NO: 8846	0.01 %	1	
10	188	SEQ ID NO: 8847	0.01 %	0.75	
11	211	SEQ ID NO: 8848	0.01 %	0.625	
12	26	SEQ ID NO: 8849	0.00 %	0.5	
13	94	SEQ ID NO: 8850	0.00 %	0.5	
14	135	SEQ ID NO: 8851	0.00 %	0.5	
15	168	SEQ ID NO: 8852	0.00 %	0.5	

	HLA A3 - 9 mers				
Maximu	m possible score us	sing this molecule type		12150	
Rank	Start position	Sequence	% of max. score	Score	
1	43	SEQ ID NO: 8853	0.24 %	30	
2	90	SEQ ID NO: 8854	0.14 %	18	
3	148	SEQ ID NO: 8855	0.09 %	12	
4	4	SEQ ID NO: 8856	0.05 %	6.75	
5	24	SEQ ID NO: 8857	0.04 %	- 6	
6	19	SEQ ID NO: 8858	0.04 %	5.4	
7	136	SEQ ID NO: 8859	0.04 %	5.4	

8	54	SEQ ID NO: 8860	0.03 %	4.5
				4.5
9	32	SEQ ID NO: 8861	0.03 %	4
10	14	SEQ ID NO: 8862	0.02 %	3.6
. 11	59	SEQ ID NO: 8863	0.02 %	3.6
12	88	SEQ ID NO: 8864	0.02 %	3
13	87	SEQ ID NO: 8865	0.02 %	2.7
14	29	SEQ ID NO: 8866	0.01 %	1.8
15	48	SEQ ID NO: 8867	0.01 %	1.8
16	115	SEQ ID NO: 8868	0.01 %	1.8
17	186	SEQ ID NO: 8869	0.01 %	1.8
18	106	SEQ ID NO: 8870	0.01 %	1.5
19	53	SEQ ID NO: 8871	0.01 %	1.35
_ 20	173	SEQ ID NO: 8872	0.00 %	1.2

HLA A3 - 10 mers					
		sing this molecule type		12150	
Rank	Start position	Sequence	% of max. score	Score	
1	24	SEQ ID NO: 8873	0.22 %	27	
2	54	SEQ ID NO: 8874	0.18 %	22.5	
3	135	SEQ ID NO: 8875	0.08 %	10.8	
4	51	SEQ ID NO: 8876	0.07 %	9	
5	13	SEQ ID NO: 8877	0.06 %	8.1	
6	26	SEQ ID NO: 8878	0.04 %	6	
_ 7	31	SEQ ID NO: 8879	0.04 %	6	
8	90	SEQ ID NO: 8880	0.04 %	6	
9	43	SEQ ID NO: 8881	0.03 %	4.5	
10	19	SEQ ID NO: 8882	0.03 %	4.05	
11	169	SEQ ID NO: 8883	0.02 %	3	
12	87	SEQ ID NO: 8884	0.02 %	2.7	
13	84	SEQ ID NO: 8885	0.01 %	1.8	
14	88	SEQ ID NO: 8886	0.01 %	1.8	
15	94	SEQ ID NO: 8887	0.01 %	1.8	
16	64	SEQ ID NO: 8888	0.00 %	1.2	
17	131	SEQ ID NO: 8889	0.00 %	1.2	
18	99	SEQ ID NO: 8890	0.00 %	1	
19	53	SEQ ID NO: 8891	0.00 %	0.9	
20	85	SEQ ID NO: 8892	0.00 %	0.9	

HLA A24 - 9 mers

Maximu	1596.672			
Rank	Start position	Sequence	% of max. score	Score
1	196	SEQ ID NO: 8893	27.55 %	440
2	44	SEQ ID NO: 8894	18.78 %	300
3	36	SEQ ID NO: 8895	12.52 %	200
4	92	SEQ ID NO: 8896	12.52 %	200
5	109	SEQ ID NO: 8897	2.70 %	43.2
6	25	SEQ ID NO: 8898	1.87 %	30
7	93	SEQ ID NO: 8899	1.12 %	18
8	12	SEQ ID NO: 8900	0.75 %	12
9	123	SEQ ID NO: 8901	0.70 %	11.2
10	7	SEQ ID NO: 8902	0.64 %	10.368
11	17	SEQ ID NO: 8903	0.52 %	8.4
12	139	SEQ ID NO: 8904	0.52 %	8.4
13	193	SEQ ID NO: 8905	0.46 %	7.5
14	6	SEQ ID NO: 8906	0.45 %	7.2
15	19	SEQ ID NO: 8907	0.45 %	7.2
16	110	SEQ ID NO: 8908	0.45 %	7.2
17	114 '	SEQ ID NO: 8909	0.45 %	7.2
18	210	SEQ ID NO: 8910	0.45 %	7.2
19	46	SEQ ID NO: 8911	0.42 %	6.72
20	52	SEQ ID NO: 8912	0.37 %	6

HLA A24 - 10 mers				
Maximu	Maximum possible score using this molecule type			
Rank	Start position	Sequence	% of max. score	Score
1	92	SEQ ID NO: 8913	7.51 %	120
2	42	SEQ ID NO: 8914	2.63 %	42
3	109	SEQ ID NO: 8915	2.25 %	36
4	23	SEQ ID NO: 8916	1.87 %	30
5	34	SEQ ID NO: 8917	0.75 %	12
6	6	SEQ ID NO: 8918	0.64 %	10.368
7	45	SEQ ID NO: 8919	0.63 %	10.08
8	196	SEQ ID NO: 8920	0.62 %	10
9	44	SEQ ID NO: 8921	0.56 %	9
10	40	SEQ ID NO: 8922	0.55 %	8.8
11	62	SEQ ID NO: 8923	0.46 %	7.5
12	193	SEQ ID NO: 8924	0.46 %	7.5
13	18	SEQ ID NO: 8925	0.45 %	7.2

14	113	SEQ ID NO: 8926	0.45 %	7.2
15	56	SEQ ID NO: 8927	0.37 %	6
16	176	SEQ ID NO: 8928	0.37 %	6
17	16	SEQ ID NO: 8929	0.35 %	5.6
18	138	SEQ ID NO: 8930	0.35 %	5.6
19	127	SEQ ID NO: 8931	0.33 %	5.28
20	36	SEQ ID NO: 8932	0.31 %	5

	HLA A 0201 - 9 mers				
Maxim	Maximum possible score using this molecule type 3925227.1				
Rank	Start position	Sequence	% of max. score	Score	
1	13	SEQ ID NO: 8933	0.04 %	1793.676528	
2	87	SEQ ID NO: 8934	0.03 %	1415.3832	
3	24	SEQ ID NO: 8935	0.01 %	618.0996816	
4	19	SEQ ID NO: 8936	0.00 %	223.23708	
5	12	SEQ ID NO: 8937	0.00 %	210.36400875	
6	. 51	SEQ ID NO: 8938	0.00 %	198.30859992	
7	53	SEQ ID NO: 8939	0.00 %	194.477328	
8	88	SEQ ID NO: 8940	0.00 %	180.58536756	
9	106	SEQ ID NO: 8941	0.00 %	169.74828	
10	54	SEQ ID NO: 8942	0.00 %	70.09848	
11	59	SEQ ID NO: 8943	0.00 %	43.42032	
12	94	SEQ ID NO: 8944	0.00 %	41.792058	
13	20	SEQ ID NO: 8945	0.00 %	37.46088108	
14	63	SEQ ID NO: 8946	0.00 %	35.73520902	
15	22	SEQ ID NO: 8947	0.00 %	20.5916435109	
16	47	SEQ ID NO: 8948	0.00 %	12.233222865	
17	66	SEQ ID NO: 8949	0.00 %	12.2199	
18	56	SEQ ID NO: 8950	0.00 %	11.486706	
19	67	SEQ ID NO: 8951	0.00 %	6.416172	
20	117	SEQ ID NO: 8952	0.00 %	5.827464	

	HLA A 0201 - 10 mers				
Maxim	um possible scor	3925227.1			
Rank	k Start position Sequence % of max. score		Score		
1	43	SEQ ID NO: 8953		3977.8497792	
2	24	SEQ ID NO: 8954	0.02 %	836.2525104	
3	51	SEQ ID NO: 8955	0.02 %	815.616432	
_ 4	49	SEQ ID NO: 8956	0.01 %	660.3245145	

5	19	SEQ ID NO: 8957	0.00 %	251.837856
6	59	SEQ ID NO: 8958	0.00 %	159.9696
7	12	SEQ ID NO: 8959	0.00 %	155.245377
8	45	SEQ ID NO: 8960	0.00 %	141.1974531
9	21	SEQ ID NO: 8961	0.00 %	117.22672269
10	53	SEQ ID NO: 8962	0.00 %	84.55536
11	87	SEQ ID NO: 8963	0.00 %	65.5671672
12	13	SEQ ID NO: 8964	0.00 %	64.88888616
13	153	SEQ ID NO: 8965	0.00 %	49.13352
14	178	SEQ ID NO: 8966	0.00 %	26.082
15	18	SEQ ID NO: 8967	0.00 %	24.802259691
16	116	SEQ ID NO: 8968	0.00 %	21.5616168
17	65	SEQ ID NO: 8969	0.00 %	20.77383
18	86	SEQ ID NO: 8970	0.00 %	15.7068219
19	27	SEQ ID NO: 8971	0.00 %	12.3159135
20	· 46	SEQ ID NO: 8972	0.00 %	11.45624789925

	HLA A 1101 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	4	SEQ ID NO: 8973	12.5 %	4.5	
2	136	SEQ ID NO: 8974	3.33 %	1.2	
3	156	SEQ ID NO: 8975	3.33 %	1.2	
4	140	SEQ ID NO: 8976	1.66 %	0.6	

	HLA A 1101 - 10 mers					
Maximum possible score using this molecule type						
Rank	Start position	Sequence	% of max. score	Score		
1	169	SEQ ID NO: 8977	5.55 %	2		
2	94	SEQ ID NO: 8978	3.33 %	1.2		

	HLA B7 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	146	SEQ ID NO: 8979	0.74 %	40	
2	154	SEQ ID NO: 8980	0.74 %	40	
3	80	SEQ ID NO: 8981	0.66 %	36	
4	139	SEQ ID NO: 8982	0.33 %	18	
5	83_	SEQ ID NO: 8983	0.22 %	12	

_	200			
6	209	SEQ ID NO: 8984	0.22 %	12
7	7	SEQ ID NO: 8985	0.11 %	6
8	3	SEQ ID NO: 8986	0.07 %	4
9	6	SEQ ID NO: 8987	0.07 %	4
10	12	SEQ ID NO: 8988	0.07 %	4
11	19	SEQ ID NO: 8989	0.07 %	4
12	24	SEQ ID NO: 8990	0.07 %	4
13	38	SEQ ID NO: 8991	0.07 %	- 4
14	46	SEQ ID NO: 8992	0.07 %	4
15	56	SEQ ID NO: 8993	0.07 %	4
16	110	SEQ ID NO: 8994	0.07 %	4
17	114	SEQ ID NO: 8995	0.07 %	4
18	123	SEQ ID NO: 8996	0.07 %	4
19	129	SEQ ID NO: 8997	0.07 %	4
.20	166	SEQ ID NO: 8998	0.07 %	4

	HLA B7 - 10 mers			
Maximu		sing this molecule type		5400
Rank		Sequence	% of max. score	Score
1	56	SEQ ID NO: 8999	1.48 %	80
2	40	SEQ ID NO: 9000	0.74 %	40
3	127	SEQ ID NO: 9001	0.74 %	40
4	170	SEQ ID NO: 9002	0.74 %	40
5	140	SEQ ID NO: 9003	0.27 %	15
6	35	SEQ ID NO: 9004	0.22 %	12
7	79	SEQ ID NO: 9005	0.22 %	12
8	82	SEQ ID NO: 9006	0.22 %	12
9	208	SEQ ID NO: 9007	0.22 %	12
10	209	SEQ ID NO: 9008	0.22 %	12
11	80	SEQ ID NO: 9009	0.16 %	9
12	129	SEQ ID NO: 9010	0.14 %	8
13	138	SEQ ID NO: 9011	0.11 %	6
14	73 .	SEQ ID NO: 9012	0.09 %	5
15	2	SEQ ID NO: 9013	0.07 %	4
16	5	SEQ ID NO: 9014	0.07 %	4
17	6	SEQ ID NO: 9015	0.07 %	4
18	16	SEQ ID NO: 9016	0.07 %	4
19	18	SEQ ID NO: 9017	0.07 %	4
20	24	SEQ ID NO: 9018	0.07 %	4

	HLA A1 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	53	SEQ ID NO: 9019	2 %	112.5	
2	10	SEQ ID NO: 9020	0.08 %	⁻ 4.5	
3	33	SEQ ID NO: 9021	0.02 %	1.5	
4	3	SEQ ID NO: 9022	0.00 %	0.5	
5	27	SEQ ID NO: 9023	0.00 %	0.5	
6	29	SEQ ID NO: 9024	0.00 %	0.5	

	HLA A1 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	10	SEQ ID NO: 9025	0.8 %	45	
2	52	SEQ ID NO: 9026	0.2 %	11.25	
3	50	SEQ ID NO: 9027	0.04 %	2.5	
4	32	SEQ ID NO: 9028	0.02 %	1.5	
5	48	SEQ ID NO: 9029	0.02 %	1.35	
6	27	SEQ ID NO: 9030	0.00 %	0.5	

	HLA A3 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	38	SEQ ID NO: 9031	1.85 %	225	
2	17	SEQ ID NO: 9032	0.02 %	3.6	
3	2	SEQ ID NO: 9033	0.02 %	2.7	
4	37	SEQ ID NO: 9034	0.01 %	1.8	
5	27	SEQ ID NO: 9035	0.01 %	1.35	
6	13	SEQ ID NO: 9036	0.00 %	0.675	
7	14	SEQ ID NO: 9037	0.00 %	0.6	

HLA A3 - 10 mers				
Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score
1	13	SEQ ID NO: 9038	0.04 %	6
2	37	SEQ ID NO: 9039	0.01 %	2.025

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3	2	SEQ ID NO: 9040	0.00 %	0.9
4	19	SEQ ID NO: 9041	0.00 %	0.675
5	16	SEQ ID NO: 9042	0.00 %	0.54

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HLA A24 - 9 mers				
Maximu	ım possible score ı	using this molecule ty	pe	1596.672
Rank	Start position	Sequence	% of max. score	Score
1	20	SEQ ID NO: 9043	1.25 %	-20
2	6	SEQ ID NO: 9044	0.52 %	8.4
3	5	SEQ ID NO: 9045	0.51 %	8.25
4	35	SEQ ID NO: 9046	0.36 %	5.76
5	31 `	SEQ ID NO: 9047	0.35 %	5.6
6	43	SEQ ID NO: 9048	0.27 %	4.4
7	13	SEQ ID NO: 9049	0.26 %	4.2
8	32	SEQ ID NO: 9050	0.21 %	3.36
9	2	SEQ ID NO: 9051	0.11 %	1.8
10	9	SEQ ID NO: 9052	0.10 %	1.68
11	8	SEQ ID NO: 9053	0.09 %	1.5
12	15	SEQ ID NO: 9054	0.09 %	1.5
13	23	SEQ ID NO: 9055	0.09 %	1.5
14	27	SEQ ID NO: 9056	0.08 %	1.4
15	24	SEQ ID NO: 9057	0.07 %	1.2
16	. 7	SEQ ID NO: 9058	0.06 %	1
17	17	SEQ ID NO: 9059	0.06 %	1
18	10	SEQ ID NO: 9060	0.05 %	0.9
19	39	SEQ ID NO: 9061	0.04 %	0.792
20	47	SEQ ID NO: 9062	0.04 %	0.792

	HLA A24 - 10 mers				
Maximu	ım possible score ı	using this molecule ty	ре	1596.672	
Rank	Start position	Sequence	% of max. score	Score	
1	5	SEQ ID NO: 9063	2.63 %	42	
2	34	SEQ ID NO: 9064	0.54 %	8.64	
3	30	SEQ ID NO: 9065	0.52 %	8.4	
4	19	SEQ ID NO: 9066	0.50 %	8	
5	50	SEQ ID NO: 9067	0.33 %	5.28	
6	12	SEQ ID NO: 9068	0.26 %	4.2	
7	31	SEQ ID NO: 9069	0.21 %	3.36	
8	26	SEQ ID NO: 9070	0.15 %	2.52	

9	8	SEQ ID NO: 9071	0.13 %	2.1
10	22	SEQ ID NO: 9072	0.12 %	2
11	23	SEQ ID NO: 9073	0.11 %	1.8
12	6	SEQ ID NO: 9074	0.09 %	1.5
13	14	SEQ ID NO: 9075	0.09 %	1.5
14	16	SEQ ID NO: 9076	0.09 %	1.5
15	7	SEQ ID NO: 9077	0.06 %	1
16	48	SEQ ID NO: 9078	0.04 %	0.75
17	0	SEQ ID NO: 9079	0.04 %	0.72
18	9	SEQ ID NO: 9080	0.04 %	0.72
19	47	SEQ ID NO: 9081	0.04 %	0.66
20	39	SEQ ID NO: 9082	0.03 % `	0.6

	HLA A 0201 - 9 mers				
Maxim	um possible score	using this molecule	e type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	15	SEQ ID NO: 9083	0.00 %	14.1442686	
2	27	SEQ ID NO: 9084	0.00 %	9.598176	
3	22	SEQ ID NO: 9085	0.00 %	9.5634	
4	9	SEQ ID NO: 9086	0.00 %	5.546246013	
5	2	SEQ ID NO: 9087	0.00 %	5.526462816	
6	24	SEQ ID NO: 9088	0.00 %	4.88163753	
7	17	SEQ ID NO: 9089	0.00 %	3.699285408	
8	31	SEQ ID NO: 9090	0.00 %	2.29699206	
9	6	SEQ ID NO: 9091	0.00 %	2.0016040674	
10	. 7	SEQ ID NO: 9092	0.00 %	0.91287	
_ 11_	49	SEQ ID NO: 9093	0.00 %	0.71805678	
12	16	SEQ ID NO: 9094	0.00 %	0.6694257042	
13	12	SEQ ID NO: 9095	0.00 %	0.6539828625	

	HLA A 0201 - 10 mers					
	Maximum possible score using this molecule type					
Rank	Start position	Sequence	% of max. score	Score		
1	16	SEQ ID NO: 9096	0.00 %	34.28765802		
2	19	SEQ ID NO: 9097	0.00 %	18.9368775		
3	14	SEQ ID NO: 9098	0.00 %	14.1442686		
4	27	SEQ ID NO: 9099	0.00 %	11.406528		
_ 5	26	SEQ ID NO: 9100	0.00 %	10.9304361558		
6	34	SEQ ID NO: 9101	0.00 %	5.580927		

7	6	SEQ ID NO: 9102	0.00 %	4.865742
8	9	SEQ ID NO: 9103	0.00 %	2.64106953
9	50	SEQ ID NO: 9104	0.00 %	2.6275752
10	30	SEQ ID NO: 9105	0.00 %	2.29699206
11	7	SEQ ID NO: 9106	0.00 %	0.86083641
12	42	SEQ ID NO: 9107	0.00 %	0.7049592
13	22	SEQ ID NO: 9108	0.00 %	0.6628440357
14	2	SEQ ID NO: 9109	0.00 %	0.6530644656

HLA A 1101 - 9 mers				
Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score
1	37	SEQ ID NO: 9110	15 %	5.4
2	38	SEQ ID NO: 9111	2.22 %	0.8

HLA A 1101 - 10 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	37	SEQ ID NO: 9112	7.5 %	2.7	

	HLA B7 - 9 mers				
Maximu		sing this molecule type		5400	
Rank	Start position	Sequence	% of max. score	Score	
11	35	SEQ ID NO: 9113	3.70 %	200	
2	17	SEQ ID NO: 9114	0.11 %	6	
3	6	SEQ ID NO: 9115	0.07 %	4	
. 4	20	SEQ ID NO: 9116	0.07 %	4	
5	31	SEQ ID NO: 9117	0.07 %	4	
6	43	SEQ ID NO: 9118	0.07 %	4	
.7	7	SEQ ID NO: 9119	0.03 %	2	
8	23	SEQ ID NO: 9120	0.02 %	1.2	
9	24	SEQ ID NO: 9121	0.02 %	1.2	
10	10	SEQ ID NO: 9122	0.01 %	0.9	

HLA B7 - 10 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	35	SEQ ID NO: 9123	0.09 %	5	
2	19	SEQ ID NO: 9124	0.07 %	4	

3	30	SEQ ID NO: 9125	0.07 %	4
4	34	SEQ ID NO: 9126	0.07 %	4
5	7	SEQ ID NO: 9127	0.03 %	2
6	16	SEQ ID NO: 9128	0.03 %	1.8
7	23	SEQ ID NO: 9129	0.02 %	1.2
8	50	SEQ ID NO: 9130	0.02 %	1.2
9	9	SEQ ID NO: 9131	0.01 %	1

Table 22: Epitopes for SEQ ID NO: 6048

HLA A1 - 9 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	66	SEQ ID NO: 9132	0.44 %	25	
2	80 ,	SEQ ID NO: 9133	0.08 %	5	
3	- 93	SEQ ID NO: 9134	0.04 %	2.7	
4	11	SEQ ID NO: 9135	0.04 %	2.5	
5	89	SEQ ID NO: 9136	0.04 %	2.25	
6	48	SEQ ID NO: 9137	0.01 %	1	
7	3	SEQ ID NO: 9138	0.00 %	0.5	
8	9	SEQ ID NO: 9139	0.00 %	0.5	
9	56	SEQ ID NO: 9140	0.00 %	0.5	
10	101	SEQ ID NO: 9141	0.00 %	0.5	
11	106	SEQ ID NO: 9142	0.00 %	0.5	
12	110	SEQ ID NO: 9143	0.00 %	0.5	

	HLA A1 - 10 mers				
Maximu	m possible score us	sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1	30	SEQ ID NO: 9144	0.4 %	22.5	
2	88	SEQ ID NO: 9145	0.12 %	6.75	
3	48	SEQ ID NO: 9146	0.04 %	2.5	
4	55	SEQ ID NO: 9147	0.02 %	1.25	
5	13	SEQ ID NO: 9148	0.01 %	0.9	
6	79	SEQ ID NO: 9149	0.01 %	0.75	
7	93	SEQ ID NO: 9150	0.01 %	0.675	
8	2	SEQ ID NO: 9151	0.00 %	0.5	
9	8	SEQ ID NO: 9152	0.00 %	0.5	

		The second secon		
10	65	SEQ ID NO: 9153	0.00 %	0.5
11	66	SEQ ID NO: 9154	0.00 %	0.5
12	80	SEQ ID NO: 9155	0.00 %	0.5
13	105	SEQ ID NO: 9156	0.00 %	0.5
14	109	SEQ ID NO: 9157	0.00 %	0.5

HLA A3 - 9 mers				
Maximu	m possible score us	sing this molecule type		12150
Rank	Start position	Sequence	% of max. score	Score
1	109	SEQ ID NO: 9158	0.74 %	90
_2	3	SEQ ID NO: 9159	0.24 %	30
3	111	SEQ ID NO: 9160	0.12 %	15
4	106	SEQ ID NO: 9161	0.07 %	9
5	95	SEQ ID NO: 9162	0.05 %	6.075
-6	101	SEQ ID NO: 9163	0.04 %	6
7 ·	110	SEQ ID NO: 9164	0.02 %	3.6
- 8	84	SEQ ID NO: 9165	0.02 %	3
9	· 80	SEQ ID NO: 9166	0.02 %	2.7
10	37	SEQ ID NO: 9167	0.01 %	2.25
11	9	SEQ ID NO: 9168	0.01 %	2
12	54	SEQ ID NO: 9169	0.01 %	2
13	99	SEQ ID NO: 9170	0.01 %	1.35
14	1	SEQ ID NO: 9171	0.01 %	1.215
15	11	SEQ ID NO: 9172	0.00 %	0.9
16	15	SEQ ID NO: 9173	0.00 %	0.9
17	69	SEQ ID NO: 9174	0.00 %	0.6
18	5	SEQ ID NO: 9175	0.00 %	0.54
19	103	SEQ ID NO: 9176	0.00 %	0.54

	HLA A3 - 10 mers				
	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	75	SEQ ID NO: 9177	0.49 %	60	
_ 2	109	SEQ ID NO: 9178	0.29 %	36	
3	22	SEQ ID NO: 9179	0.14 %	18	
4	15	SEQ ID NO: 9180	0.04 %	6	
5	110	SEQ ID NO: 9181	0.01 %	2.25	
6	95	SEQ ID NO: 9182	0.01 %	1.8	
7	101	SEQ ID NO: 9183	0.01 %	1.35	

8	43	SEQ ID NO: 9184	0.00 %	1
9	2	SEQ ID NO: 9185	0.00 %	0.9
10	5	SEQ ID NO: 9186	0.00 %	0.9
11	ア	SEQ ID NO: 9187	0.00 %	0.9
12	107	SEQ ID NO: 9188	0.00 %	0.9
13	102	SEQ ID NO: 9189	0.00 %	0.81
14	3	SEQ ID NO: 9190	0.00 %	0.75
15	8	SEQ ID NO: 9191	0.00 %	-0.6
16	103	SEQ ID NO: 9192	0.00 %	0.54

	HLA A24 - 9 mers				
Maximu	ım poṡsible score ι	using this molecule ty	pe	1596.672	
Rank	Start position	Sequence	% of max. score	Score	
1	88	SEQ ID NO: 9193	1.66 %	26.6112	
2	77	SEQ ID NO: 9194	0.77 %	12.32	
3	18	SEQ ID NO: 9195	0.56 %	9	
4	108	SEQ ID NO: 9196	0.56 %	9	
• 5	92	SEQ ID NO: 9197	0.54 %	8.64	
6	96	SEQ ID NO: 9198	0.54 %	8.64	
7	73	SEQ ID NO: 9199	0.46 %	7.5	
8	40	SEQ ID NO: 9200	0.45 %	7.2	
9	104	SEQ ID NO: 9201	0.42 %	6.72	
10	. 8	SEQ ID NO: 9202	0.41 %	6.6	
11	21	SEQ ID NO: 9203	0.37 %	6	
12	102	SEQ ID NO: 9204	0.37 %	6	
13	22	SEQ ID NO: 9205	0.25 %	4	
14	68	SEQ ID NO: 9206	0.25 %	4	
15	106	SEQ ID NO: 9207	0.22 %	3.6	
16	1	SEQ ID NO: 9208	0.18 %	3	
17	79	SEQ ID NO: 9209	0.18 %	3	
18	93	SEQ ID NO: 9210	0.18 %	3	
19	101	SEQ ID NO: 9211	0.18 %	3	
20	37	SEQ ID NO: 9212	0.15 %	2.4	

	HLA A24 - 10 mers					
Maximu	Maximum possible score using this molecule type					
Rank	Start position	Sequence	% of max. score	Score		
1	100	SEQ ID NO: 9213	0.93 %	15		
2	18	SEQ ID NO: 9214	0.78 %	12.6		

3	98	SEQ ID NO: 9215	0.52 %	8.4
4	73	SEQ ID NO: 9216	0.46 %	7.5
5	91	SEQ ID NO: 9217	0.45 %	7.2
6	103	SEQ ID NO: 9218	0.42 %	6.72
7	7	SEQ ID NO: 9219	0.41 %	6.6
8	21	SEQ ID NO: 9220	0.37 %	6
9	46	SEQ ID NO: 9221	0.37 %	6
10	93	SEQ ID NO: 9222	0.37 %	-6
11	96	SEQ ID NO: 9223	0.37 %	6
12	101	SEQ ID NO: 9224	0.37 %	6
13	77	SEQ ID NO: 9225	0.25 %	4
14	92	SEQ ID NO: 9226	0.22 %	3.6
15	105	SEQ ID NO: 9227	0.22 %	3.6
16	2	SEQ ID NO: 9228	0.18 %	3
17	53	SEQ ID NO: 9229	0.18 %	3
18	36	SEQ ID NO: 9230	0.12 %	2
19	55	SEQ ID NO: 9231	0.12 %	2
20	102	SEQ ID NO: 9232	0.11 %	1.8

	HLA A 0201 - 9 mers				
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	84	SEQ ID NO: 9233	0.01 %	441.342216	
2	102	SEQ ID NO: 9234	0.00 %	63.16728165	
3	107	SEQ ID NO: 9235	0.00 %	51.882640425	
4	11	SEQ ID NO: 9236	0.00 %	43.8816609	
5	95	SEQ ID NO: 9237	0.00 %	33.40165248	
6	2	SEQ ID NO: 9238	0.00 %	24.66305226	
7	92	SEQ ID NO: 9239	0.00 %	22.64458905	
8	103	SEQ ID NO: 9240	0.00 %	20.70206586	
9	47	SEQ ID NO: 9241	0.00 %	11.175953184	
10	94	SEQ ID NO: 9242	0.00 %	8.452983	
11	15	SEQ ID NO: 9243	0.00 %	8.1793152	
12	8	SEQ ID NO: 9244	0.00 %	4.993461	
13	5	SEQ ID NO: 9245	0.00 %	4.57284528	
14	99	SEQ ID NO: 9246	0.00 %	3.999468528	
_15	105	SEQ ID NO: 9247	0.00 %	2.231322	
16	20	SEQ ID NO: 9248	0.00 %	1.3524	
17	62	SEQ ID NO: 9249	0.00 %	0.8631693	

18	6	SEQ ID NO: 9250	0.00 %	0.824619
19	57	SEQ ID NO: 9251	0.00 %	0.72105
20	58	SEQ ID NO: 9252	0.00 %	0.7147572

	HLA A 0201 - 10 mers				
Maxim	um possible score	using this molecule	type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	101	SEQ ID NO: 9253	0.03 %	1243.078056	
2	3	SEQ ID NO: 9254	0.01 %	592.944462	
3	106	SEQ ID NO: 9255	0.00 %	94.2678	
4	5	SEQ ID NO: 9256	0.00 %	43.42032	
5	107	SEQ ID NO: 9257	0.00 %	33.30332334	
6	102	SEQ ID NO: 9258	0.00 %	32.53181778	
7	54	SEQ ID NO: 9259	0.00 %	27.324	
8	7	SEQ ID NO: 9260	0.00 %	21.3624	
9	1	SEQ ID NO: 9261	0.00 %	13.723479	
10	95	SEQ ID NO: 9262	0.00 %	13.00344192	
11	94	SEQ ID NO: 9263	0.00 %	10.01276388	
12	99	SEQ ID NO: 9264	0.00 %	5.6615328	
13	39	SEQ ID NO: 9265	0.00 %	3.6304212	
14	111	SEQ ID NO: 9266	0.00 %	2.53368	
15	103	SEQ ID NO: 9267	0.00 %	2.475394803	
16	14	SEQ ID NO: 9268	0.00 %	2.4519012	
17	19	SEQ ID NO: 9269	0.00 %	2.07604992	
18	29	SEQ ID NO: 9270	0.00 %	1.8179154	
19	57	SEQ ID NO: 9271	0.00 %	1.52076	
20	47	SEQ ID NO: 9272	0.00 %	1.27712376	

	HLA A 1101 - 9 mers					
Maximum possible score using this molecule type						
Rank	Rank Start position Sequence % of max. score					
1	80	SEQ ID NO: 9273	3.33 %	1.2		
2	69	SEQ ID NO: 9274	1.66 %	0.6		
3	109	SEQ ID NO: 9275	1.66 %	0.6		

	HLA A 1101 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	22	SEQ ID NO: 9276	11.11 %	4	

HLA B7 - 9 mers				
		sing this molecule type		5400
Rank	Start position	Sequence	% of max. score	Score
11	22	SEQ ID NO: 9277	3.70 %	200
2	77	SEQ ID NO: 9278	2.22 %	120
3	104	SEQ ID NO: 9279	0.22 %	12
4	40	SEQ ID NO: 9280	0.11 %	- 6
5	8	SEQ ID NO: 9281	0.07 %	4
6	21	SEQ ID NO: 9282	0.07 %	4
7	68	SEQ ID NO: 9283	0.07 %	4
8	92	SEQ ID NO: 9284	0.07 %	4
9	102	SEQ ID NO: 9285	0.07 %	4
10	46	SEQ ID NO: 9286	0.03 %	2
11	98	SEQ ID NO: 9287	0.03 %	2
12	103	SEQ ID NO: 9288	0.03 %	2
13	88	SEQ ID NO: 9289	0.02 %	1.2
14	105	SEQ ID NO: 9290	0.01 %	0.9
15	43	SEQ ID NO: 9291	0.01 %	0.6
16	79	SEQ ID NO: 9292	0.01 %	0.6
17	95	SEQ ID NO: 9293	0.01 %	0.6
18	107	SEQ ID NO: 9294	0.00 %	0.5

HLA B7 - 10 mers						
Maximu	Maximum possible score using this molecule type 5400					
Rank	Start position	Sequence	% of max. score	Score		
11	46	SEQ ID NO: 9295	1.48 %	80		
2	98	SEQ ID NO: 9296	1.48 %	80		
3	91	SEQ ID NO: 9297	0.37 %	20		
4	103	SEQ ID NO: 9298	0.37 %	20		
5	7	SEQ ID NO: 9299	0.07 %	4		
_ 6	21	SEQ ID NO: 9300	0.07 %	4		
7	101	SEQ ID NO: 9301	0.07 %	4		
8	107	SEQ ID NO: 9302	0.03 %	2		
9	67	SEQ ID NO: 9303	0.02 %	1.2		
10	93	SEQ ID NO: 9304	0.02 %	1.2		
11	69	SEQ ID NO: 9305	0.01 %	1		
12	39	SEQ ID NO: 9306	0.01 %	0.6		
13	77	SEQ ID NO: 9307	0.01 %	0.6		

SEO ID NO: 9308		
	I 0.00 % I	

Table 23: Epitopes for SEQ ID NO: 6049

	HLA A1 - 9 mers				
Maximu	m possible score us	sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1	0	SEQ ID NO: 9309	0.2 %	11.25	
2	35	SEQ ID NO: 9310	0.01 %	0.9	
3	4	SEQ ID NO: 9311	0.00 %	0.5	
4	5	SEQ ID NO: 9312	0.00 %	0.5	
5	10	SEQ ID NO: 9313	0.00 %	0.5	
6	19	SEQ ID NO: 9314	0.00 %	0.5	
7	21	SEQ ID NO: 9315	0.00 %	0.5	

	HLA A1 - 10 mers					
Maximu	Maximum possible score using this molecule type					
Rank	Rank Start position Sequence % of max. score					
1	0	SEQ ID NO: 9316	0.2 %	11.25		
2	5	SEQ ID NO: 9317	0.04 %	2.5		
3	33	SEQ ID NO: 9318	0.02 %	1.5		
4	3	SEQ ID NO: 9319	0.02 %	1.25		
5	9 .	SEQ ID NO: 9320	0.00 %	0.5		
6	18	SEQ ID NO: 9321	0.00 %	0.5		
7	20	SEQ ID NO: 9322	0.00 %	0.5		

	HLA A3 - 9 mers					
Maximu	faximum possible score using this molecule type 12150					
Rank	Start position	Sequence	Sequence % of max. score			
1	4	SEQ ID NO: 9323	0.14 %	18		
2	16	SEQ ID NO: 9324	0.11 %	13.5		
3	23	SEQ ID NO: 9325	0.06 %	8.1		
4	18	SEQ ID NO: 9326	0.03 %	4.05		
5	21	SEQ ID NO: 9327	0.01 %	2.025		
6	9	SEQ ID NO: 9328	0.01 %	1.8		
7	15	SEQ ID NO: 9329	0.01 %	1.8		
8	25	SEQ ID NO: 9330	0.01 %	1.8		
9	12	SEQ ID NO: 9331	0.00 %	0.9		

10	19	SEQ ID NO: 9332	0.00 %	0.9
11	20	SEQ ID NO: 9333	0.00 %	0.9
12	2	SEQ ID NO: 9334	0.00 %	0.81
13	22	SEQ ID NO: 9335	0.00 %	0.81
14	10	SEQ ID NO: 9336	0.00 %	0.6

L	HLA A3 - 10 mers					
Maximu	Maximum possible score using this molecule type					
Rank	Start position	Sequence	Sequence % of max. score			
11	20	SEQ ID NO: 9337	0.16 %	20.25		
2	9	SEQ ID NO: 9338	0.09 %	12		
3	16	SEQ ID NO: 9339	0.07 %	9		
4	18	SEQ ID NO: 9340	0.07 %	9		
5	22	SEQ ID NO: 9341	0.06 %	8.1		
6	4	SEQ ID NO: 9342	0.03 %	4.05		
. 7	15	SEQ ID NO: 9343	0.03 %	4.05		
8	12	SEQ ID NO: 9344	0.02 %	3.6		
9	3	SEQ ID NO: 9345	0.00 %	0.9		
10	33	SEQ ID NO: 9346	0.00 %	0.6		
11	· 2	SEQ ID NO: 9347	0.00 %	0.54		
12	24	SEQ ID NO: 9348	0.00 %	0.54		

	HLA A24 - 9 mers					
	Maximum possible score using this molecule type					
Rank	Start position	Sequence	% of max. score	Score		
1	8	SEQ ID NO: 9349	18.78 %	300		
2	11	SEQ ID NO: 9350	1.87 %	30		
3	28	SEQ ID NO: 9351	1.50 %	24		
4	7	SEQ ID NO: 9352	0.75 %	12		
5	17	SEQ ID NO: 9353	0.56 %	9		
6	14	SEQ ID NO: 9354	0.46 %	7.5		
7	23	SEQ ID NO: 9355	0.37 %	6		
8	13	SEQ ID NO: 9356	0.36 %	5.76		
9	· 2	SEQ ID NO: 9357	0.35 %	5.6		
10	16	SEQ ID NO: 9358	0.35 %	5.6		
11	9	SEQ ID NO: 9359	0.30 %	4.8		
12	21	SEQ ID NO: 9360	0.26 %	4.2		
13	5	SEQ ID NO: 9361	0.25 %	4		
14	4	SEQ ID NO: 9362	0.22 %	3.6		

15	0	SEQ ID NO: 9363	0.18 %	3
16	19	SEQ ID NO: 9364	0.18 %	3
17	10	SEQ ID NO: 9365	0.15 %	2.4
18	18	SEQ ID NO: 9366	0.13 %	2.1
19	25	SEQ ID NO: 9367	0.06 %	1.1
20	15	SEQ ID NO: 9368	0.05 %	0.9

	HLA A24 - 10 mers					
Maximu	ım possible score ı	using this molecule ty	ре	1596.672		
Rank	Start position	Sequence	% of max. score	Score		
1	8	SEQ ID NO: 9369	22.54 %	360		
2	7	SEQ ID NO: 9370	1.25 %	. 20		
3	17	SEQ ID NO: 9371	0.65 %	10.5		
4	15	SEQ ID NO: 9372	0.52 %	8.4		
5	4	SEQ ID NO: 9373	0.45 %	7.2		
6	22	SEQ ID NO: 9374	0.37 %	6		
7	12	SEQ ID NO: 9375	0.36 %	5.76		
. 8	27	SEQ ID NO: 9376	0.30 %	4.8		
9	14	SEQ ID NO: 9377	0.28 %	4.5		
10	20	SEQ ID NO: 9378	0.26 %	4.2		
11	10	SEQ ID NO: 9379	0.25 %	4		
12	3	SEQ ID NO: 9380	0.18 %	. 3		
13	18	SEQ ID NO: 9381	0.18 %	. 3		
14	9	SEQ ID NO: 9382	0.15 %	2.4		
15	24	SEQ ID NO: 9383	0.10 %	1.65		
16	16	SEQ ID NO: 9384	0.07 %	1.2		
17	13	SEQ ID NO: 9385	0.06 %	1		
18	11	SEQ ID NO: 9386	0.05 %	0.9		
19	1	SEQ ID NO: 9387	0.05 %	0.84		

	HLA A 0201 - 9 mers					
		e using this molecul	e type	3925227.1		
Rank	Start position	Sequence	% of max. score	Score		
1	12	SEQ ID NO: 9388	0.10 %	4267.988928		
2	23	SEQ ID NO: 9389	0.03 %	1360.69088544		
3	9	SEQ ID NO: 9390	0.01 %	569.948832		
4	16	SEQ ID NO: 9391	0.00 %	309.0498408		
5	15	SEQ ID NO: 9392	0.00 %	79.73570448		
6	2	SEQ ID NO: 9393	0.00 %	51.109542		

18	SEQ ID NO: 9394	0.00.00	
		0.00 %	45.25539984
25	SEQ ID NO: 9395	0.00 %	34.28765802
22	SEQ ID NO: 9396	0.00 %	26.532116325
5	SEQ ID NO: 9397	0.00 %	25.26691266
21	SEQ ID NO: 9398	0.00 %	4.72873208445
11	SEQ ID NO: 9399	0.00 %	2.638538265
8	SEQ ID NO: 9400	0.00 %	2,4274552038
4	SEQ ID NO: 9401	0.00 %	1.7415324
20	SEQ ID NO: 9402	0.00 %	1.6025526
13	SEQ ID NO: 9403	0.00 %	1.453803297
35	SEQ ID NO: 9404	0.00 %	1.36878336
3 .	SEQ ID NO: 9405		0.824619
33	SEQ ID NO: 9406	0.00 %	0.513774
	5 21 11 8 4 20 13 35 3	5 SEQ ID NO: 9397 21 SEQ ID NO: 9398 11 SEQ ID NO: 9399 8 SEQ ID NO: 9400 4 SEQ ID NO: 9401 20 SEQ ID NO: 9402 13 SEQ ID NO: 9403 35 SEQ ID NO: 9404 3 SEQ ID NO: 9405	22 SEQ ID NO: 9396 0.00 % 5 SEQ ID NO: 9397 0.00 % 21 SEQ ID NO: 9398 0.00 % 11 SEQ ID NO: 9399 0.00 % 8 SEQ ID NO: 9400 0.00 % 4 SEQ ID NO: 9401 0.00 % 20 SEQ ID NO: 9402 0.00 % 13 SEQ ID NO: 9403 0.00 % 35 SEQ ID NO: 9404 0.00 % 3 SEQ ID NO: 9405 0.00 %

HLA A 0201 - 10 mers				
Maxim	um possible scor	re using this molecu	ile type	3925227.1
Rank	Start position	Sequence	% of max. score	Score
1	22	SEQ ID NO: 9407	0.09 %	3636.068421648
2	• 4	SEQ ID NO: 9408	0.02 %	1107.960876
3	15	SEQ ID NO: 9409	0.02 %	836.2525104
4	16	SEQ ID NO: 9410	0.00 %	150.9313176
5	12	SEQ ID NO: 9411	0.00 %	76.55002416
6	1	SEQ ID NO: 9412	0.00 %	49.0273014
7	10	SEQ ID NO: 9413	0.00 %	42.1638414747
8	20	SEQ ID NO: 9414	0.00 %	9.29480508
9	24	SEQ ID NO: 9415	0.00 %	9.2669346
10	13	SEQ ID NO: 9416	0.00 %	7.96581954
11	21	SEQ ID NO: 9417	0.00 %	5.051306761875
12	5	SEQ ID NO: 9418	0.00 %	2.6941464
13	11	SEQ ID NO: 9419	0.00 %	2.3839914
14	34	SEQ ID NO: 9420	0.00 %	1,465422
15	2	SEQ ID NO: 9421	0.00 %	0.70794
16	9	SEQ ID NO: 9422	0.00 %	0.6513048
17	19	SEQ ID NO: 9423	0.00 %	0.51882640425

HLA A 1101 - 9 mers				
Maximum possible score using this molecule type				36
Rank Start position Sequence % of max. score Score				

HLA A 1101 - 10 mers				
Maximu	Maximum possible score using this molecule type			
Rank	Start position	Sequence	% of max. score	Score
11	33	SEQ ID NO: 9424	1.66 %	0,6

	HLA B7 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	13	SEQ ID NO: 9425	0.22 %	12	
2	2	SEQ ID NO: 9426	0.07 %	4	
3	9	SEQ ID NO: 9427	0.07 %	4	
4	16	SEQ ID NO: 9428	0.07 %	4	
5	23	SEQ ID NO: 9429	0.07 %	4	
6	5	SEQ ID NO: 9430	0.02 %	1.2	
7	15	SEQ ID NO: 9431	0.01 %	1	

	HLA B7 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Rank Start position Sequence % of max. score				
1	4	SEQ ID NO: 9432	0.07 %	4	
:2	10	SEQ ID NO: 9433	0.07 %	4	
3	12	SEQ ID NO: 9434	0.07 %	4	
4	15	SEQ ID NO: 9435	0.07 %	4	
5	22	SEQ ID NO: 9436	0.07 %	4	
6	13	SEQ ID NO: 9437	0.02 %	1.2	

5 Table 24: Epitopes for SEQ ID NO: 6050

HLA A1 - 9 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	47	SEQ ID NO: 9438	0.01 %	0.75	
2	21	SEQ ID NO: 9439	0.00 %	0.5	
3	53	SEQ ID NO: 9440	0.00 %	0.5	

HLA A1 - 10 mers				
Maximu	Maximum possible score using this molecule type			5625
				Score

1	16	SEQ ID NO: 9441	0.04 %	2.5
2	71	SEQ ID NO: 9442	0.04 %	2.5
3	47	SEQ ID NO: 9443	0.02 %	1.5
4	62	SEQ ID NO: 9444	0.01 %	0.9
5	20	SEQ ID NO: 9445	0.00 %	0.5
6	38	SEQ ID NO: 9446	0.00 %	0.5

	HLA A3 - 9 mers				
Maximum possible score using this molecule type					
Rank	Start position	Sequence	% of max. score	Score	
1	54	SEQ ID NO: 9447	0.02 %	2.7	
2	17	SEQ ID NO: 9448	0.01 %	2	
3	3	SEQ ID NO: 9449	0.01 %	1.8	

	HLA A3 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
11	22 .	SEQ ID NO: 9450	0.09 %	12	
2	16	SEQ ID NO: 9451	0.01 %	2	
3	54	SEQ ID NO: 9452	0.00 %	0.9	

	HLA A24 - 9 mers				
		using this molecule ty	pe	1596.672	
Rank	Start position	Sequence	% of max. score	Score	
1	70	SEQ ID NO: 9453	2.10 %	33.6	
2	77	SEQ ID NO: 9454	1.12 %	18	
3	60	SEQ ID NO: 9455	0.46 %	7.5	
4	54	SEQ ID NO: 9456	0.37 %	6	
5	14 ,	SEQ ID NO: 9457	0.31 %	5	
6	19	SEQ ID NO: 9458	0.30 %	4.8	
7	47	SEQ ID NO: 9459	0.30 %	4.8	
8	12	SEQ ID NO: 9460	0.25 %	4	
9	15	SEQ ID NO: 9461	0.25 %	4	
10	67	SEQ ID NO: 9462	0.25 %	4	
11	21	SEQ ID NO: 9463	0.18 %	3	
12	· 37	SEQ ID NO: 9464	0.06 %	1	
13	27	SEQ ID NO: 9465	0.03 %	0.5	

HLA A24 - 10 mers

Maximu	ım possible score ı	using this molecule ty	pe	1596.672
Rank	Start position	Sequence	% of max. score	Score
1	14	SEQ ID NO: 9466	12.52 %	200
2	7	SEQ ID NO: 9467	0.93 %	15
3	11	SEQ ID NO: 9468	0.75 %	12
4	60	SEQ ID NO: 9469	0.56 %	9
5	18	SEQ ID NO: 9470	0.45 %	7.2
6	46	SEQ ID NO: 9471	0.45 %	7.2
7	53	SEQ ID NO: 9472	0.37 %	6
8	69	SEQ ID NO: 9473	0.35 %	5.6
9	66	SEQ ID NO: 9474	0.25 %	4
10	20	SEQ ID NO: 9475	0.12 %	2
11	47	SEQ ID NO: 9476	0.07 %	1.2
12	36	SEQ ID NO: 9477	0.06 %	1
13	26	SEQ ID NO: 9478	0.04 %	0.75
14	70	SEQ ID NO: 9479	0.04 %	0.72

HLA A 0201 - 9 mers						
Maximu	3925227.1					
Rank	Start position	Sequence	% of max. score	Score		
1	54	SEQ ID NO: 9480	0.02 %	881.199		
2	26	SEQ ID NO: 9481	0.00 %	95.013		
3	61	SEQ ID NO: 9482	0.00 %	93.69648		
4	19	SEQ ID NO: 9483	0.00 %	40.2894864		
5	74	SEQ ID NO: 9484	0.00 %	12.6684		
6	35	SEQ ID NO: 9485	0.00 %	10.34586		
7	69	SEQ ID NO: 9486	0.00 %	3.3704706		
8	13	SEQ ID NO: 9487	0.00 %	1.656		
9	15	SEQ ID NO: 9488	0.00 %	1.47537042		
10	68	SEQ ID NO: 9489	0.00 %	0.966		
11	22	SEQ ID NO: 9490	0.00 %	0.942678		
12	12	SEQ ID NO: 9491	0.00 %	0.7669695		
13	36	SEQ ID NO: 9492	0.00 %	0.52661835		

HLA A 0201 - 10 mers							
Maxim	3925227.1						
Rank	Start position	Sequence	% of max. score	Score			
1	61	SEQ ID NO: 9493	0.00 %	93.69648			
2	25	SEQ ID NO: 9494	0.00 %	63.33035625			

34	SEQ ID NO: 9495	0.00 %	50,232
53	SEQ ID NO: 9496	0.00 %	45.2838375
26	SEQ ID NO: 9497		14.35752
27	SEQ ID NO: 9498	0.00 %	2.8557858
17	SEQ ID NO: 9499		2.3973222
36	SEQ ID NO: 9500		1.798209
69	SEQ ID NO: 9501		1.03521597
67	SEQ ID NO: 9502	0.00 %	0.966
68	SEQ ID NO: 9503	0.00 %	0.910938
11	SEQ ID NO: 9504	0.00 %	0.7669695
	53 26 27 17 36 69	53 SEQ ID NO: 9496 26 SEQ ID NO: 9497 27 SEQ ID NO: 9498 17 SEQ ID NO: 9499 36 SEQ ID NO: 9500 69 SEQ ID NO: 9501 67 SEQ ID NO: 9502 68 SEQ ID NO: 9503	53 SEQ ID NO: 9496 0.00 % 26 SEQ ID NO: 9497 0.00 % 27 SEQ ID NO: 9498 0.00 % 17 SEQ ID NO: 9499 0.00 % 36 SEQ ID NO: 9500 0.00 % 69 SEQ ID NO: 9501 0.00 % 67 SEQ ID NO: 9502 0.00 % 68 SEQ ID NO: 9503 0.00 %

HLA A 1101 - 9 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	17	SEQ ID NO: 9505	2.22 %	0.8	

HLA A 1101 - 10 mers				
Rank	Maximum possible score using this molecule type			36
Kank	Start position	Sequence	% of max. score	Score
	16	SEQ ID NO: 9506	5.55 %	2

	HLA B7 - 9 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	5400	
11	27	SEQ ID NO: 9507	0.37 %	Score	
. 2	54	SEQ ID NO: 9508	0.22 %	20	
3	70	SEQ ID NO: 9509	0.22 %	12	
4	67	SEQ ID NO: 9510		12	
5	12	SEQ ID NO: 9511	0.11 %	6	
6	15	SEQ ID NO: 9512	0.07 %	4	
7	19	SEQ ID NO: 9513	0.07 %	4	
8	49	SEQ ID NO: 9514	0.07 %	4	
9	69	SEQ ID NO: 9515	0.03 %	2	
10	47	SEQ ID NO: 9516	0.03 %	1.8	
11	5	SEQ ID NO: 9516	0.02 %	1.2	
12	9		0.01 %	1	
13	35	SEQ ID NO: 9518	0.01 %	11	
14	37	SEQ ID NO: 9519	0.01 %	1	
15		SEQ ID NO: 9520	0.01 %	0.6	
	68	SEQ ID NO: 9521	0.01 %	0.6	

	HLA B7 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	5400 Score	
1	69	SEQ ID NO: 9522	0.66 %	36	
2	53	SEQ ID NO: 9523	0.22 %	12	
3	5	SEQ ID NO: 9524	0.13 %	7.5	
4	66	SEQ ID NO: 9525	0.11 %	6	
5	11	SEQ ID NO: 9526	0.07 %	4	
6	27	SEQ ID NO: 9527	0.07 %	4	
7	46	SEQ ID NO: 9528	0.07 %	4	
8	18 ·	SEQ ID NO: 9529	0.02 %	1.2	
9	9	SEQ ID NO: 9530	0.01 %	1	
10	26	SEQ ID NO: 9531	0.01 %	1	
11	25	SEQ ID NO: 9532	0.01 %	0.75	
12	17	SEQ ID NO: 9533	0.01 %	0.6	
13	36	SEQ ID NO: 9534	0.01 %	0.6	
14	68	SEQ ID NO: 9535	0.01 %	0.6	
15	35	SEQ ID NO: 9536	- 0.00 %	0.5	
16	42	SEQ ID NO: 9537	0.00 %	0.5	
17	73	SEQ ID NO: 9538	0.00 %	0.5	

Table 25: Epitopes for SEQ ID NO: 6052

	HLA A1 - 9 mers				
Maximu	m possible score u	sing this molecule type		5625	
Rank	Start position	Sequence	% of max. score	Score	
1_1_	365	SEQ ID NO: 9539	0.8 %	45	
2	397	SEQ ID NO: 9540	0.44 %	25	
3	229	SEQ ID NO: 9541	0.32 %	18	
4	103	· SEQ ID NO: 9542	0.17 %	10	
5	338	SEQ ID NO: 9543	0.17 %	10	
6	251	SEQ ID NO: 9544	0.16 %	9	
7	79	SEQ ID NO: 9545	0.11 %	6.25	
8	119	SEQ ID NO: 9546	0.10 %	6	
9	361	SEQ ID NO: 9547	0.08 %	5	
10	60	SEQ ID NO: 9548	0.04 %	2,25	
11	101	SEQ ID NO: 9549	0.04 %	2.25	
12	278	SEQ ID NO: 9550	0.04 %	2.25	

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13	23	SEQ ID NO: 9551	0.02 %	1.25
_14	164	SEQ ID NO: 9552	0.02 %	1.25
15	165	SEQ ID NO: 9553	0.02 %	1.25
16	295	SEQ ID NO: 9554	0.02 %	1.25
17	172	SEQ ID NO: 9555	0.01 %	0.9
18	0	SEQ ID NO: 9556	0.01 %	0.75
19	311	SEQ ID NO: 9557	0.01 %	0.75
20	78	SEQ ID NO: 9558	0.01 %	0.625

	HLA A1 - 10 mers			
Maximu	Maximum possible score using this molecule type			
Rank	Start position	Seguence	% of max. score	Score
11	114	SEQ ID NO: 9559	1.11 %	62.5
2	134	SEQ ID NO: 9560	0.8 %	45
3	365	SEQ ID NO: 9561	0.8 %	45
4	77	SEQ ID NO: 9562	0.66 %	37.5
5	. 103	SEQ ID NO: 9563	0.44 %	25
6	23 -	SEQ ID NO: 9564	0.22 %	12.5
7 *	. 338	SEQ ID NO: 9565	0.17 %	1.0
8	361	SEQ ID NO: 9566	0.17 %	10
9	324	SEQ ID NO: 9567	0.11 %	6.25
10	375	SEQ ID NO: 9568	0.11 %	6.25
11	79	SEQ ID NO: 9569	0.04 %	2.5
12	295	SEQ ID NO: 9570	0.04 %	2.5
13	346	SEQ ID NO: 9571	0.04 %	2.5
14	378	SEQ ID NO: 9572	0.03 %	2
15	251	SEQ ID NO: 9573	0.03 %	1.8
16	214	SEQ ID NO: 9574	0.02 %	1.125
17	160	SEQ ID NO: 9575	0.01 %	1
18	172	SEQ ID NO: 9576	0.01 %	0.9
19	229	SEQ ID NO: 9577	0.01 %	0.9
20	376	SEQ ID NO: 9578	0.01 %	0.9

	HLA A3 - 9 mers				
Maximum possible score using this molecule type				12150	
Rank	Start position	Sequence	% of max. score	Score	
1	229	SEQ ID NO: 9579	0.49 %	60	
2	361	SEQ ID NO: 9580	0.27 %	33.75	
3	330	SEQ ID NO: 9581	0.16 %	20	

4	218	SEQ ID NO: 9582	0.09 %	12
5	338	SEQ ID NO: 9583	0.04 %	6
6	352	SEQ ID NO: 9584	0.04 %	6 .
7	103	SEQ ID NO: 9585	0.04 %	5.4
8	291	SEQ ID NO: 9586	0.01 %	2
9	241	SEQ ID NO: 9587	0.01 %	1.8
10	290	SEQ ID NO: 9588	0.01 %	1.8
11	316	SEQ ID NO: 9589	0.01 %	1.8
12	222	SEQ ID NO: 9590	0.01 %	1.35
13	266	SEQ ID NO: 9591	0.01 %	1.35
14	53	SEQ ID NO: 9592	0.00 %	1
15	100	SEQ ID NO: 9593	0.00 %	0.9
16	138	SEQ ID NO: 9594	0.00 %	0.9
17	240	SEQ ID NO: 9595	0.00 %	0.9
18	119	SEQ ID NO: 9596	0.00 %	0.675
19	44	SEQ ID NO: 9597	0.00 %	0.6
20	161	SEQ ID NO: 9598	0.00 %	0.6

HLA A3 - 10 mers				
Maximu	Maximum possible score using this molecule type			
Rank	Start position	Sequence	% of max. score	Score
1	338	SEQ ID NO: 9599	0.49 %	60
2	160	SEQ ID NO: 9600	0.32 %	40
3	352	SEQ ID NO: 9601	0.24 %	30
4	361	SEQ ID NO: 9602	0.18 %	22.5
5	103	SEQ ID NO: 9603	0.13 %	16.2
6	290	SEQ ID NO: 9604	0.07 %	9
7	351	SEQ ID NO: 9605	0.07 %	9
8	44	SEQ ID NO: 9606	0.04 %	6
9	228	SEQ ID NO: 9607	0.03 %	4.05
10	394	SEQ ID NO: 9608	0.02 %	3
11	240	SEQ ID NO: 9609	0.02 %	2.7
12	100	SEQ ID NO: 9610	0.01 %	1.8
13	114	SEQ ID NO: 9611	0.01 %	1.8
14	93	SEQ ID NO: 9612	0.01 %	1.5
15	134	SEQ ID NO: 9613	0.01 %	1.5
16	221	SEQ ID NO: 9614	0.01 %	1.35
17	330	SEQ ID NO: 9615	0.00 %	1.2
18	112	SEQ ID NO: 9616	0.00 %	0.9

10	240			
19	218	SEQ ID NO: 9617	0.00 %	0.0
20	EE	CEO ID NO COLO	0.00 /0	0.9
		SEQ ID NO: 9618	0.00 %	0.6

	HLA A24 - 9 mers				
Maxim	um possible score	using this molecule ty	/ne	1506 670	
Rank	Start position	Sequence	% of max. score	1596.672	
1	345	SEQ ID NO: 9619	1.50 %	Score	
2	306	SEQ ID NO: 9620		24	
3	222	SEQ ID NO: 9621	0.75 %	12	
4	111	SEQ ID NO: 9622	0.54 %	8.64	
5	159		0.51 %	8.25	
6		SEQ ID NO: 9623	0.45 %	7.2	
7	219	SEQ ID NO: 9624	0.45 %	7.2	
	283	SEQ ID NO: 9625	0.45 %	7.2	
8	266	SEQ ID NO: 9626	0.42 %	6.72	
9	56	SEQ ID NO: 9627	0.41 %	6.6	
10	131	SEQ ID NO: 9628	0.37 %	6	
11	214	SEQ ID NO: 9629	0.37 %		
12	297	SEQ ID NO: 9630	0.37 %	6	
13	86	SEQ ID NO: 9631	0.31 %	5	
14	122	SEQ ID NO: 9632	0.31 %	5	
15	48	SEQ ID NO: 9633	0.30 %	4.8	
16	105	SEQ ID NO: 9634	0.30 %	4.8	
17	213	SEQ ID NO: 9635	0.30 %		
18	323	SEQ ID NO: 9636	0.30 %	4.8	
19	338	SEQ ID NO: 9637	0.30 %		
20	399	SEQ ID NO: 9638	0.30 %	4.8	

96.672 Score
15
12
10.56
9.6
8
7.5
7.2
7.2
1.2

10	277	SEQ ID NO: 9648	0.45 %	7.2
11	150	SEQ ID NO: 9649	0.37 %	6
12	152	SEQ ID NO: 9650	0.37 %	6
13	158	SEQ ID NO: 9651	0.37 %	6
14	171	SEQ ID NO: 9652	0.37 %	6
15	343	SEQ ID NO: 9653	0.37 %	6
16	110	SEQ ID NO: 9654	0.34 %	5.5
17	85	SEQ ID NO: 9655	0.31 %	5
18	47	SEQ ID NO: 9656	0.30 %	4.8
19	213	SEQ ID NO: 9657	0.30 %	4.8
20	218	SEQ ID NO: 9658	0.30 %	4.8

	HLA A 0201 - 9 mers				
Maxim	um possible scor	e using this molecul	e type	3925227.1	
Rank	Start position	Sequence	% of max. score	Score	
1	222	SEQ ID NO: 9659	0.03 %	1267.10434728	
2	226	SEQ ID NO: 9660	0.00 %	69.552	
3	316	SEQ ID NO: 9661	0.00 %	50.232	
4	351	SEQ ID NO: 9662	0.00 %	31.24872	
5	159	SEQ ID NO: 9663	0.00 %	13.6235739	
6	406	SEQ ID NO: 9664	0.00 %	11.4264	
7	165	SEQ ID NO: 9665	0.00 %	8.14407	
8	238	SEQ ID NO: 9666	0.00 %	7.0518	
9	138	SEQ ID NO: 9667	0.00 %	5.112072	
10	130	SEQ ID NO: 9668	0.00 %	3.00547233	
11	303	SEQ ID NO: 9669	0.00 %	2.59578	
12	157	SEQ ID NO: 9670	0.00 %	2.412585	
13	219	SEQ ID NO: 9671	0.00 %	2.103255861	
14	305	SEQ ID NO: 9672	0.00 %	1.86369	
15	158	SEQ ID NO: 9673	0.00 %	1.646892	
16	331	SEQ ID NO: 9674	0.00 %	1.614048	
17	399	SEQ ID NO: 9675	0.00 %	1.442246832	
18	324	SEQ ID NO: 9676	0.00 %	1.319625	
19	312	SEQ ID NO: 9677	0.00 %	1.233099	
20	262	SEQ ID NO: 9678	0.00 %	0.966	

HLA A 0201 - 10 mers				
Maximum possible score using this molecule type			3925227.1	
Rank Start position Sequence % of max. score Score				

11	221	SEQ ID NO: 9679	0.00 %	309.0498408
_ 2	112	SEQ ID NO: 9680	0.00 %	98.26704
3	330	SEQ ID NO: 9681	0.00 %	98.26704
4	158	SEQ ID NO: 9682	0.00 %	36.31608
5	218	SEQ ID NO: 9683	0.00 %	24.0754248
6_	124	SEQ ID NO: 9684	0.00 %	12.2199
_ 7	. 55	SEQ ID NO: 9685	0.00 %	10.467576
8	315	SEQ ID NO: 9686	0.00 %	7.7274204
9	350	SEQ ID NO: 9687	0.00 %	4.296699
10	405	SEQ ID NO: 9688	0.00 %	4.286487
11	388	SEQ ID NO: 9689	0.00 %	4.054785
12	322	SEQ ID NO: 9690	0.00 %	3.883803
13	130	SEQ ID NO: 9691	0.00 %	3.428691903
14	45	SEQ ID NO: 9692	0.00 %	3.411230625
15	132	SEQ ID NO: 9693	0.00 %	2.99943
16	410	SEQ ID NO: 9694	0.00 %	2.63718
17	. 316	SEQ ID NO: 9695	0.00 %	2.48686074
18	104	SEQ ID NO: 9696	0.00 %	2.477311485
19	164	SEQ ID NO: 9697	0.00 %	2.2011
20	282	SEQ ID NO: 9698	0.00 %	2.16591

HLA A 1101 - 9 mers					
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	361	SEQ ID NO: 9699	16.66 %	6	
. 2	53	SEQ ID NO: 9700	2.77 %	1	
3	240	SEQ ID NO: 9701	1.66 %	0.6	
4	241	SEQ ID NO: 9702	1.66 %	0.6	

	HLA A 1101 - 10 mers				
	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	361	SEQ ID NO: 9703	16.66 %	6	
2	93	SEQ ID NO: 9704	8.33 %	3	
3	338	SEQ ID NO: 9705	3.33 %	1.2	
4	134	SEQ ID NO: 9706	2.77 %	1	
_ 5	228	SEQ ID NO: 9707	2.5 %	0.9	
6	160	SEQ ID NO: 9708	2.22 %	0.8	
7	239	SEQ ID NO: 9709	1.66 %	0.6	

8	240	SEQ ID NO: 9710	1.66 %	0.6
9	257	SEQ ID NO: 9711	1.66 %	0.6
10	379	SEO ID NO: 9712	1.66 %	0.6

	HLA B7 - 9 mers				
Maximu	m possible score us	sing this molecule type		5400	
Rank	Start position	Sequence	% of max. score	Score	
1	105	SEQ ID NO: 9713	14.81 %	-800	
2	66	SEQ ID NO: 9714	1.48 %	80	
3	93	SEQ ID NO: 9715	0.92 %	50	
4	257	SEQ ID NO: 9716	0.55 %	30	
5	323	SEQ ID NO: 9717	0.37 %	. 20	
6	211	SEQ ID NO: 9718	0.22 %	12	
7	219	SEQ ID NO: 9719	0.22 %	. 12	
8	. 403	SEQ ID NO: 9720	0.18 %	10	
9	343	SEQ ID NO: 9721	0.14 %	8	
10	12	SEQ ID NO: 9722	0.11 %	6	
11	113	SEQ ID NO: 9723	0.11 %	. 6	
12	48	SEQ ID NO: 9724	0.07 %	4	
13	56	SEQ ID NO: 9725	0.07 %	4	
14	150	SEQ ID NO: 9726	0.07 %	4	
15	153	SEQ ID NO: 9727	0.07 %	4	
16	159	SEQ ID NO: 9728	0.07 %	4	
17	213	SEQ ID NO: 9729	0.07 %	4.	
18	216	SEQ ID NO: 9730	0.07 %	4	
19	222	SEQ ID NO: 9731	0.07 %	4	
20	283	SEQ ID NO: 9732	0.07 %	4	

	HLA B7 - 10 mers				
Maximu	Maximum possible score using this molecule type				
Rank	Start position	Sequence	% of max. score	Score	
1	36	SEQ ID NO: 9733	1.48 %	80	
2	150	SEQ ID NO: 9734	1.48 %	80	
3	343	SEQ ID NO: 9735	1.48 %	80	
4	12	SEQ ID NO: 9736	1.11 %	60	
5	308	SEQ ID NO: 9737	1.11 %	60	
6	130	SEQ ID NO: 9738	0.37 %	20	
7	55	SEQ ID NO: 9739	0.22 %	12	
8	210	SEQ ID NO: 9740	0.22 %	12	

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9 218 SEQ ID NO: 9741 0.22 % 12 10 201 SEQ ID NO: 9742 0.18 % 10 11 121 SEQ ID NO: 9743 0.14 % 8 12 391 SEQ ID NO: 9744 0.13 % 7.5 13 112 SEQ ID NO: 9745 0.11 % 6 14 385 SEQ ID NO: 9746 0.11 % 6 15 47 SEQ ID NO: 9747 0.07 % 4 16 66 SEQ ID NO: 9748 0.07 % 4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4 20 158 SEQ ID NO: 9752 0.07 % 4					
11 121 SEQ ID NO: 9743 0.14 % 8 12 391 SEQ ID NO: 9744 0.13 % 7.5 13 112 SEQ ID NO: 9745 0.11 % 6 14 385 SEQ ID NO: 9746 0.11 % 6 15 47 SEQ ID NO: 9747 0.07 % 4 16 66 SEQ ID NO: 9748 0.07 % -4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	9	218	SEQ ID NO: 9741	0.22 %	12
12 391 SEQ ID NO: 9744 0.13 % 7.5 13 112 SEQ ID NO: 9745 0.11 % 6 14 385 SEQ ID NO: 9746 0.11 % 6 15 47 SEQ ID NO: 9747 0.07 % 4 16 66 SEQ ID NO: 9748 0.07 % -4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	10	201	SEQ ID NO: 9742	0.18 %	10
13 112 SEQ ID NO: 9745 0.11 % 6 14 385 SEQ ID NO: 9746 0.11 % 6 15 47 SEQ ID NO: 9747 0.07 % 4 16 66 SEQ ID NO: 9748 0.07 % -4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	11	121	SEQ ID NO: 9743	0.14 %	8
14 385 SEQ ID NO: 9746 0.11 % 6 15 47 SEQ ID NO: 9747 0.07 % 4 16 66 SEQ ID NO: 9748 0.07 % -4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	12	391	SEQ ID NO: 9744	0.13 %	7.5
15 47 SEQ ID NO: 9747 0.07 % 4 16 66 SEQ ID NO: 9748 0.07 % -4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	13	112	SEQ ID NO: 9745	0.11 %	6
16 66 SEQ ID NO: 9748 0.07 % - 4 17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	14	385	SEQ ID NO: 9746	0.11 %	6
17 95 SEQ ID NO: 9749 0.07 % 4 18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	15	47 -	SEQ ID NO: 9747	0.07 %	4
18 104 SEQ ID NO: 9750 0.07 % 4 19 152 SEQ ID NO: 9751 0.07 % 4	16	66	SEQ ID NO: 9748	0.07 %	- 4
19 152 SEQ ID NO: 9751 0.07 % 4	17	95	SEQ ID NO: 9749	0.07 %	4
3.07.70	18	104	SEQ ID NO: 9750	0.07 %	4
20 158 SEQ ID NO: 9752 0.07 % 4	19	152	SEQ ID NO: 9751	0.07 %	4
	20	158	SEQ ID NO: 9752	0.07 %	4

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TABLE 26: Cloned sequences for E.coli expression

ORF	DNA length	Cloning	
	bp	pET	pGEX
P28	537	NdeI / XhoI	
P65	1917	NheI / HindIII	
Nsp1A	2495	NheI / XhoI	
Nsp1B	2153	NdeI / XhoI	
Nsp1C	2612	NdeI / XhoI	
Nsp2A	431	NdeI / XhoI	BamHI / XhoI
Nsp2B	426	NdeI / XhoI	BamHI / XhoI
Nsp3	870	NdeI / XhoI	
Nsp4	249	NdeI / XhoI	BamHI / XhoI
Nsp5	594	NheI / XhoI	
Nsp6	339	NdeI / XhoI	BamHI / XhoI
Nsp7	417	NdeI / XhoI	BamHI / XhoI
Nsp9A	1385	Nhel /Xhol	
Nsp9B	1409	NdeI / XhoI	
Nsp10	1803	NheI / XhoI	
Nsp11	1581	NdeI / XhoI	
Nsp12	1038	NdeI / HindIII	
Nsp13	897	NdeI / XhoI	-
Spike (S1)	1946	NdeI / XhoI	
Spike (S2)	1598	NdeI / XhoI	
Spike (S1-S2)	3545	NdeI / XhoI	
HR1	287	NdeI / XhoI	BamHI / XhoI
HR2	146	NdeI / XhoI	BamHI / XhoI
ORF3∆100	525	NdeI / XhoI	
ORF4	465	NdeI / XhoI	
Envelope (E)	231	NdeI / XhoI	BamHI / XhoI
Matrix (M)Δ100	366	NdeI / XhoI	BamHI / XhoI
ORF7Δ18	137	NdeI / XhoI	BamHI / XhoI
ORF8	369	NdeI / XhoI	BamHI / XhoI
ORF9	135	Ndel / Xhol	BamHI / XhoI
ORF10	120	NheI / XhoI	BamHI / XhoI
ORF11	255	NdeI / XhoI	BamHI / XhoI
Nucleocapsid (N)	1269	NdeI / EcoRI	
ORF12	297	NdeI / EcoRI	BamHI / EcoRI

TABLE 27: Primers

ORF	Forward primer	Reverse primer
P28	9803	9818
P65	9804	9819
Nsp1A	9805	9820
Nsp1B	9806	9821
Nsp1C	9807	9822
Nsp2 + Nsp3	9808	9823
Nsp4 to Nsp7	9809	9824
Nsp9A	9810	9825
Nsp9B	9811	9826
Nsp10	9812	9827
Nsp11	9813	9828
Nsp12-Nsp13	9814	9829
ORF3-ORF4	9815	9830
Env-ORF10	9816	. 9831
ORF11-ORF12	9817	9832

TABLE 28: Primers

ORF	Forward primer	Reverse primer
Nsp2A	SEQ ID NO: 9833	SEQ ID NO: 9858
Nsp2B	SEQ ID NO: 9834	SEQ ID NO: 9859
Nsp3	SEQ ID NO: 9835	SEQ ID NO: 9860
Nsp4	SEQ ID NO: 9836	SEQ ID NO: 9861
Nsp5	SEQ ID NO: 9837	SEQ ID NO: 9862
Nsp6	SEQ ID NO: 9838	SEQ ID NO: 9863
Nsp7	SEQ ID NO: 9839	SEQ ID NO: 9864
Nsp12	SEQ ID NO: 9840	SEQ ID NO: 9865
Nsp13	SEQ ID NO: 9841	SEQ ID NO: 9866
Spike S1	SEQ ID NO: 9842	SEQ ID NO: 9867
Spike S2	SEQ ID NO: 9843	SEO ID NO: 9868
Spike S1-S2	SEQ ID NO: 9844	SEQ ID NO: 9869
HR1	SEQ ID NO: 9845	SEQ ID NO: 9870
HR2	SEQ ID NO: 9846	SEQ ID NO: 9871
· Orf3∆100	SEQ ID NO: 9847	SEO ID NO: 9872
Orf4	SEQ ID NO: 9848	SEQ ID NO: 9873
Env E	SEQ ID NO: 9849	SEQ ID NO: 9874
Matrix MΔ100	SEQ ID NO: 9850	SEQ ID NO: 9875
Orf7∆18	SEQ ID NO: 9851	SEO ID NO: 9876
Orf8	SEQ ID NO: 9852	SEQ ID NO: 9877
Orf9	SEQ ID NO: 9853	SEQ ID NO: 9878
Orf10	SEQ ID NO: 9854	SEO ID NO: 9879
Orf11	SEQ ID NO: 9855	SEO ID NO: 9880
Nucleocapsid N	SEQ ID NO: 9856	SEO ID NO: 9881
Orf12	SEQ ID NO: 9857	SEO ID NO: 9882

TABLE 29: Cloning, purification and expression in E.coli

SARS CoV ORFs	M.W Kd	cloning	Expr.	purification as
P28	19,7			his sol
P65	70,3	±	±	his sol
Nsp1A(N-term)	91,6	±	±	his ins
Nsp1B (core)	80,8		-	
Nsp1C (C-term)	95.3		-	
Nsp2A (N-term)	15,8	t	±	his ins
Nsp2B (C-term)	15,5	±	±	his sol
Nsp3	31,9	±		
Nsp4	9,1	+		his sol
Nsp5	21,8	+	+	his sol
Nsp6	12.4	±	±	his sol
Nsp7	15,3	+	±	his ins
Nsp9A (N-term)	50,8	+		
Nsp9B (C-term)	51,6	+		his ins
Nsp10	66			
Nsp11	58	-		
Nsp12	38			
Nsp13	32.7	<u>+</u>	+	his ins
Spike (S1-his)	71,3			his ins
Spike (S2-his)	58,6	±		
Spike (S1S2-his)	130	±	+	his ins
HR1	11		±	his ins
HR2	5,4		+	his sol
ORF3 Δ100 ¹	19.1			
ORF4	16,9	+	+	his ins (trimer)
Envelope (E)	34,3	±	±	gst ins (IB)
Matrix (M)∆100	13.3	±	L±	his ins
ORF7∆18 ²	31	+	+	gst sol
ORF8	39,5	+	+	gst ins (IB)
ORF9	30,8	+	+	gst sol
ORF10	30.3	+_	±	gst ins (IB)
ORF11	35,2	+	L±	gst ins (IB)
Nucleocapsid (N)	43,6	+	±	his ins
ORF12	36,7	+	+	his ins

TABLE 30: E.coli expression, purity and yield

Protein	Tag	Purity (%)	Yield (mg/l)
Nsp2A (N-term)	His	95	1.7
Nsp2B (C-term)	His	95	4.1
Nsp4	His	95	12.6
Nsp5	His	95	5.88
Nsp6	His	95	8.1
P28	His	95	1
P65	His	80	0.553
HR2	His	95	11.9
HR1	His	80	2.64
Nsp1A	His	95	0.267
Spike S1-S2	His	80 .	0.381
Matrix M	His	85	12.4
ORF7	GST	85	4.9

TABLE 31: Primers

SEQ ID NO:	Rank	Model	Local	(Position)
10235	F1	1	1	(106)
10236	F2	2	1	(728)
10237	F3	3	1	(112)
10238	F4	5	2	(1331)
10239	F5	6	1	(12)
10240	F6	6	1	(346)
10241	F7	8	1	(904)
10242	F8	9	1	(1016)
10243	F9	9	1	(1015)
10244	F10	9	1	(719)
10245	F11	9	1	(720)
10246	F12	10	1	(724)
10247	R1	2	1	(1283)
10248	R2	4	1	(756)
10249	R3	4	1	(758)
10250	R4	5 .	2	(259)
10251	R5	6	1	(54)
10252	R6	7	1	(648)
10253	R7	8	1	(948)
10254	R8	8	1	(260)
10255	R9	9	1	(1282)
10256	R10	9	1	(950)
10257	R11	9	1	(756)
10258	R12	10	1	(132)

TABLE 32: Primers

Primers	List: (f	forward)		
Rank	Model	Local	Sequence	(Position)
F1	7.	1	SEQ ID NO: 10352	(290)
F2	7	ī	SEO ID NO: 10353	(291)
F3	7	ī	SEO ID NO: 10354	(294)
F4	7	ī	SEO ID NO: 10355	(292)
F5	ź	i	SEQ ID NO: 10356	(293)
F6	9	ī	SEO ID NO: 10357	(198)
F7	9	i	SEQ ID NO: 10358	(199)
		1	SEQ ID NO: 10359	(33)
F8	10	1	SEQ ID NO: 10359 SEQ ID NO: 10360	(200)
F9	11	1		(299)
F10	11		SEQ ID NO: 10361	(298)
F11	12	1	SEQ ID NO: 10362	
F12	12	1	SEQ ID NO: 10363	(297)
F13	14	1	SEQ ID NO: 10364	(35)
F14	14	1	SEQ ID NO: 10365	(34)
F15	16	1	SEQ ID NO: 10366	(300)
F16	17	1	SEQ ID NO: 10367	(295)
F17	17	1	SEQ ID NO: 10368	(296)
F18	17	1	SEQ ID NO: 10369	(175)
F19	17	1	SEQ ID NO: 10370	(36)
F20	20	1	SEQ ID NO: 10371	(202)
F21	20	1	SEQ ID NO: 10372	(201)
F22	28	1 .	SEQ ID NO: 10373	(204)
F23	28	1	SEQ ID NO: 10374	(203)
F24	29	1	SEQ ID NO: 10375	(269)
F25	29	1	SEQ ID NO: 10376	(268)
Primer	s List (r	everse)		
Rank	Model	Local	Sequence	(Position)
R1	7	1	SEQ ID NO: 10377	(337)
R2	9	1	SEQ ID NO: 10378	(229)
R3	11	1	SEQ ID NO: 10379	(230)
R4		-	SEQ ID NO: 10380	(338)
	11	1		
	11 12	1	SEQ ID NO: 10381	(207)
R5				(207) (338)
R5 R6	12 12	1	SEQ ID NO: 10381 SEQ ID NO: 10382	
R5 R6 R7	12 12 13	1	SEQ ID NO: 10381	(338)
R5 R6 R7 R8	12 12 13 14	1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384	(338) (231)
R5 R6 R7 R8 R9	12 12 13 14 14	1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385	(338) (231) (80)
R5 R6 R7 R8	12 12 13 14	1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384	(338) (231) (80) (232)
R5 R6 R7 R8 R9 R10 R11	12 12 13 14 14 15	1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10386	(338) (231) (80) (232) (82)
R5 R6 R7 R8 R9 R10 R11	12 12 13 14 14 15 16	1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387	(338) (231) (80) (232) (82) (340)
R5 R6 R7 R8 R9 R10 R11	12 12 13 14 14 15	1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10387 SEQ ID NO: 10387 SEQ ID NO: 10388	(338) (231) (80) (232) (82) (340) (83)
R5 R6 R7 R8 R9 R10 R11 R12 R13	12 12 13 14 14 15 16 17 17	1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10386 SEQ ID NO: 10388 SEQ ID NO: 10389	(338) (231) (80) (232) (82) (340) (83) (206)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14	12 12 13 14 14 15 16 17 17	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10387 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10390	(338) (231) (80) (232) (82) (340) (83) (206) (82)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15	12 12 13 14 14 15 16 17 17 17 17	1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10399 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10391	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16	12 12 13 14 14 15 16 17 17 17 17 17	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10387 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10391 SEQ ID NO: 10392 SEQ ID NO: 10393	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (340)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16	12 12 13 14 14 15 16 17 17 17 17 18 20 20	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10391 SEQ ID NO: 10392 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17	12 12 13 14 14 15 16 17 17 17 17 18 20 20 21	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10388 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10392 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10394 SEQ ID NO: 10395	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (340) (233) (79)
R5 R6 R7 R8 R9 R10 R112 R13 R14 R15 R16 R17 R18	12 12 13 14 14 15 16 17 17 17 17 20 20 21 22	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10399 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10394 SEQ ID NO: 10395 SEQ ID NO: 10395 SEQ ID NO: 10395 SEQ ID NO: 10395	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (340) (233) (79) (213)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17 R18	12 12 13 14 15 16 17 17 17 17 18 20 20 21 22 28	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10388 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10392 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10395 SEQ ID NO: 10396 SEQ ID NO: 10397	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (340) (233) (79) (213) (236)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17 R18 R19 R20 R21	12 12 13 14 14 15 16 17 17 17 17 20 20 21 22 22 28 29	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10399 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10395 SEQ ID NO: 10395 SEQ ID NO: 10395 SEQ ID NO: 10396 SEQ ID NO: 10397 SEQ ID NO: 10397	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (344) (233) (79) (213) (236) (317)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17 R17 R18 R19 R20 R21 R22 R23	12 12 13 14 15 16 17 17 17 17 20 20 21 22 28 29 32	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10395 SEQ ID NO: 10397 SEQ ID NO: 10397 SEQ ID NO: 103997 SEQ ID NO: 103998 SEQ ID NO: 103998	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (340) (233) (79) (213) (236) (317) (391)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17 R18 R19 R20 R21 R22 R23 R24	12 12 13 14 15 16 17 17 17 18 20 21 22 22 28 29 32 35	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10388 SEQ ID NO: 10389 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10395 SEQ ID NO: 10396 SEQ ID NO: 10397 SEQ ID NO: 10397 SEQ ID NO: 10398 SEQ ID NO: 10399	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (344) (233) (79) (213) (236) (317) (391) (557)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17 R18 R19 R20 R21 R22 R23 R24 R25	12 12 13 14 15 16 17 17 17 18 20 21 22 28 29 32 35 36	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10391 SEQ ID NO: 10391 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10394 SEQ ID NO: 10396 SEQ ID NO: 10397 SEQ ID NO: 10397 SEQ ID NO: 10399 SEQ ID NO: 10399 SEQ ID NO: 10399 SEQ ID NO: 10400	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (340) (233) (79) (213) (236) (317) (391) (57)
R5 R6 R7 R8 R9 R10 R11 R12 R13 R14 R15 R16 R17 R17 R20 R21 R22 R23 R24 R25	12 12 13 14 14 15 16 17 17 17 17 18 20 20 21 22 28 29 32 35 36 st (left part):	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	SEQ ID NO: 10381 SEQ ID NO: 10382 SEQ ID NO: 10383 SEQ ID NO: 10384 SEQ ID NO: 10384 SEQ ID NO: 10385 SEQ ID NO: 10386 SEQ ID NO: 10387 SEQ ID NO: 10387 SEQ ID NO: 10389 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10390 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10393 SEQ ID NO: 10395 SEQ ID NO: 10395 SEQ ID NO: 10395 SEQ ID NO: 10396 SEQ ID NO: 10397 SEQ ID NO: 10399 SEQ ID NO: 10400 SEQ ID NO: 10400 SEQ ID NO: 10400 SEQ ID NO: 10400 SEQ ID NO: 10401	(338) (231) (80) (232) (82) (340) (83) (206) (82) (337) (341) (344) (233) (79) (213) (236) (317) (391) (557)

TABLE 33: Primers

rimers List	(forward)	res			
Rank	Model	Local	Sequence	(Position)	
F1	1	1	SEO ID NO: 10580	(637)	
F2	2	ī	SEO ID NO: 10581	(439)	
F3	2	î	SEQ ID NO: 10582		
F4	3	ī	SEQ ID NO: 10583	(440)	
P5	4	1	SEQ ID NO: 10584	(729)	
F6	4	1	SEQ ID NO: 10585	(696)	
F7 .	4	1		(697)	
F8	5	1	SEQ ID NO: 10586	(111)	
F9	5		SEQ ID NO: 10587	(867)	
F10	5	1	SEQ ID NO: 10588	(868)	
		1	SEQ ID NO: 10589	(869)	
F11	5	1	SEQ ID NO: 10590	(640)	
F12	6	1	SEQ ID NO: 10591	(438)	
F13	6	1	SEQ ID NO: 10592	(437)	
F14	6	1	SEQ ID NO: 10593	(436)	
F15	6	1	SEQ ID NO: 10594	(732)	•
F16	6	1	SEQ ID NO: 10595	(635)	
F17	6	1	SEQ ID NO: 10596	(457)	•
F18	6	1	SEQ ID NO: 10597	(458)	
F19	6	1	SEQ ID NO: 10598	(636)	•
F20	7	1	SEQ ID NO: 10599	(854)	
F21	7	1	SEQ ID NO: 10600	(855)	
F22	7	1	SEQ ID NO: 10601	(581)	
F23	7	1	SEQ ID NO: 10602	(853)	
F24	7	1	SEQ ID NO: 10603	(342)	
F25	. 2	1	SEQ ID NO: 10604	(343)	
F26	7	1	SEQ ID NO: 10605	(112)	
F27	7	1	SEQ ID NO: 10606	(94)	
F28	7	1	SEQ ID NO: 10607	(642)	
F29	8	1	SEQ ID NO: 10608	(638)	
F30	8	1	SEQ ID NO: 10609	(639)	
F31	8	1	SEO ID NO: 10610	(730)	
F32	8	1	SEQ ID NO: 10611	(641)	
F33 -	8	1	SEQ ID NO: 10612	(731)	
F34	8	1	SEQ ID NO: 10613	(326)	•
F35	8	1	SEQ ID NO: 10614	(325)	
F36	9	ī	SEQ ID NO: 10615	(517)	
F37	9	1	SEQ ID NO: 10616	(701)	
F38	9	ī	SEQ ID NO: 10617	(208)	
F39	9	ī	SEQ ID NO: 10618	(209)	
F40	9	ī	SEQ ID NO: 10619	(702)	
F41	9	ī	SEQ ID NO: 10619	(210)	
F42	10	ī	SEQ ID NO: 10621	(634)	
F43	10	i	SEQ ID NO: 10621		
F44	10	1	SEQ ID NO: 10622 SEO ID NO: 10623	(694)	
F45	10	1	SEQ ID NO: 10623 SEQ ID NO: 10624	(693)	
F46	10	1		(728)	
F47	10	1	SEQ ID NO: 10625	(695)	
F48	11	1	SEQ ID NO: 10626	(95)	
F49	11	1	SEQ ID NO: 10627	(455)	
F49 F50			SEQ ID NO: 10628	(456)	
F 3 0	11	1	SEQ ID NO: 10629	(454)	

	Primers I	List (reverse)			
8		Sc	ores			9
	Rank	Mode1	Local	Sequence	(Position)	
	R1	1	1	SEQ ID NO: 10630	(367)	
1	R2	1	1	SEQ ID NO: 10631	(666)	
1	R3	2	1	SEQ ID NO: 10632	(464)	
1	R4	3	1	SEQ ID NO: 10633	(669)	
1	R5	3	1	SEQ ID NO: 10634	(750)	
	R6	4	1	SEQ ID NO: 10635	(720)	
	R7	4	1	SEQ ID NO: 10636	(465)	
	R8	4	1	SEQ ID NO: 10637	(370)	
1	R9	4 .	1	SEQ ID NO: 10638	(668)	
l	R10	4	1	SEQ ID NO: 10639	(135)	
١.	R11	5	1	SEQ ID NO: 10640	(901)	
	R12	5	1	SEQ ID NO: 10641	(667)	
	R13	6	1	SEQ ID NO: 10642	(609)	
	R14	6	1	SEQ ID NO: 10643	(464)	
l	R15	6	1	SEQ ID NO: 10644	(665)	
1	R16	6	1	SEO ID NO: 10645	(486)	
	R17	6	1	SEO ID NO: 10646	(356)	
}	R18	6	ī	SEQ ID NO: 10647	(758)	•
1	R19	7	ī	SEQ ID NO: 10648	(366)	
1	R20	7	ī	SEQ ID NO: 10649	(368)	•
1	R21	7	1	SEQ ID NO: 10650	(136)	
1	R22	7	ī	SEO ID NO: 10651	(675)	
1	R23	7	ī	SEO ID NO: 10652	(366)	
1	R24	7	ī	SEO ID NO: 10653	(608)	
i	R25 .	ź	i	SEQ ID NO: 10654	(884)	
1	R26	7	1	SEQ ID NO: 10655	(120)	
1	R27	8	1	SEQ ID NO: 10656	(355)	9
1	R28	8	1	SEQ ID NO: 10657	(671)	
1	R29	8	1	SEQ ID NO: 10658	(756)	*
1	R30	8	1	SEQ ID NO: 10659	(751)	·
1	R31	8	i	SEQ ID NO: 10660	(666)	
1	R32	9	i	SEO ID NO: 10661	(242)	
l	R33	9	1	SEQ ID NO: 10661	(543)	
1	R34 .	9	i	SEQ ID NO: 10663	(724)	
1	R35	9	1	SEQ ID NO: 10664	(482)	
1	R36	10	1	SEQ ID NO: 10665	(121)	
1	R37	10	i	SEQ ID NO: 10666	(662)	
1	R38	10	1	SEQ ID NO: 10667		•
1	R39	10	1	SEQ ID NO: 10667	(750) (719)	
1	R40	10	1	SEO ID NO: 10669		
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	R42	11	1	SEQ ID NO: 10671	(375)	3
1	R43	11	1	SEQ ID NO: 10672	(728)	
1	R44	11	1	SEQ ID NO: 10673	(373)	
	R45	11		SEQ ID NO: 10674	(998)	
	R46	11	1	SEQ ID NO: 10675	(486)	
1	R47	12	1	SEQ ID NO: 10676	(881)	
1	R48	12	1	SEQ ID NO: 10677	(882)	
1	R49	12	1	SEQ ID NO: 10678	(244)	
1	R50	12	1	SEQ ID NO: 10679	(1003)	

Primers List (left part): SEQ ID NO^{\$}: 10680-10974 | Primers List (right part): SEQ ID NO^{\$}: 10975-11282 |
Primers List (forward): SEQ ID NO^{\$}: 11283-11302 | Primers List (reverse): SEQ ID NO^{\$}: 11303-11322

TABLE 34

Compound #	Structure	Name	МH+
1	HC-CH, H-CH,	N-methyl-4-[(2-[[2-(1-methylethyl)phenyl]amino)-1H-benzimidazol-5-yl)oxy]pyridine-2-carboxamide	402.5
2	Hosi-ori	N-methyl-4-{[1-methyl-2-({3- [(trimethylsilyl)ethynyl]phenyl)am ino)-1H-benzimidazol-5- yl]oxy]pyridine-2-carboxamide	470.6
3		N-methyl-4-[(1-methyl-2-[[2- (phenylcarbonyl)phenyl]amino}- 1H-benzimidazol-5- yl)oxy]pyridine-2-carboxamide	478.5
4	0	4-(methyloxy)-N-[6-(methyloxy)- 1,3-benzothiazol-2-yl]-3- nitrobenzamide	360.4
5		4-({2-[(4-butylphenyl)amino]-1,3- benzothiazol-5-yl]oxy)-N- methylpyridine-2-carboxamide	433.5
6		N-methyl-4-([1-methyl-2-[(6- pyrrolidin-1-ylpyridin-3-yl)amino]- 1H-benzimidazol-5- yl]oxy)pyridine-2-carboxamide	444.5
7	O "H" ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	4-((2-[1,1'-bi(cyclohexyl)-2- ylamino]-1-methyl-1H- benzimidazol-5-yl]oxy)-N- methylpyridine-2-carboxamide	462.6
8	H _c	4-((2-[(4-chlorophenyl)amlno]-1- methyl-1H-benzimidazol-5- //]oxy)-N-1,3-thiazol-2- //pyridine-2-carboxamide	477.9

9		4-[(1-methyl-2-{[2- (methyloxy)phenyl]amino)-1H- benzlimidazol-5-yl)oxy]-N-[3- (methyloxy)propyl]pyridine-2- carboxamide	462.5
10		4-({2-[(4-ethylphenyl)amino]-1,3- benzoxazol-5-yl)oxy}-N- methylpyridine-2-carboxamide	389.4
11	t	1-[(3-fluorophenyl)carbonyl]-4- {[4- (trifluoromethyl)phenyl]methyl}pi perazine	367.4
: : 12	CH ₃ O CH ₃ O CH ₃	1-[2-(ethyloxy)phenyl]-4-([3,4,5- tris(methyloxy)phenyl]carbonyl)p liperazine	401.5
13	ST. C. C.	1-(3-chlorophenyl)-4-[[2- (ethyloxy)phenyl]carbonyl)pipera zine	345.8
14		3-((4-[(2E)-3-phenylprop-2- enylpiperazin-1-yl)carbonyl)-7- oxabicyclo[2.2.1]heptane-2- carboxylic acid	371.4
15	H ₂ C. O. CH ₃	1-[2-(methyloxy)phenyl]-4- {[3,4,5- tris(methyloxy)phenyl]carbonyl]p perazine	387.4
16		3-[(4-pyridin-2-ylpiperazin-1- yl)carbonyl]-7- oxabicyclo[2,2.1]heptane-2- carboxylic acid	332.4

17	HO STATE OF THE ST	3-pentyl-7-[(4-phenylpiperazin-1- yl)carbonyl]-2-thioxo-2,3- dihydroquinazolin-4(1H)-one	437.6
18	Ho COTH	1-[(E)-((4-[(2,4- dimethylphenyl)methyl]piperazin -1-yl]imino)methyl]naphthalen-2- ol	374.5
19		5-chloro-1-{[3- (trifluoromethyl)phenyl]methyl)- 1H-indole-2,3-dione	340.7
20	O.N. CH	1-[(4-methylphenyl)methyl]-5- nitro-1H-indole-2,3-dione	297.3
21	CH ₃ CH ₃ O CH ₃	1-methyl-6,7-bis(methyloxy)-2- {[3-(methyloxy)phenyl]carbonyl}- 1,2,3,4-tetrahydroisoquinoline	342.4
22	CH ₃	1-methyl-6,7-bis(methyloxy)-2- (naphthalen-2-ylcarbonyl)- 1,2,3,4-tetrahydroisoquinoline	362.4
23		[2-(trifluoromethyl)phenyl]methyl 3-[4-(amlnocarbonyl)phenyl]-2- cycloheptyl-1-oxo-1,2,3,4- tetrahydroisoquinoline-4- carboxylate	565.6
24	O N-SN	anthra[1,2-c][1,2,5]thiadiazole- 6,11-dione	267.3

25		benzo[b]oxanthrene-6,11-dione	265.2
26	N CONTRACTOR	ethyl 6,11-dioxo-6,11- dihydrobenzo[b]phenazine-2- carboxylate	333.3
27	H ₃ C _{-N} -CH ₃ 0=5=00	N,N-dimethyl-9,10-dioxo-9,10- dihydroanthracene-1- sulfonamide	316.3
28	HE CONTRACTOR	2-(trifluoromethyl)-3-[[3,4,5- tris(methyloxy)phenyl]carbonyl]n aphtho[2,3-b]furan-4,9-dione	461.4
29	CH ₃	2-(2-oxopropyl)-2-phenyl-1H- indene-1,3(2H)-dione	279.3
30		ethyl 4-{5-[(3- nitrophenyl)carbonyl]-1,3-dioxo- 1,3-dihydro-2H-isoindol-2- yl]benzoate	445.4
31	CI N F F	5,6-dichloro-2-[2-chloro-5- (trifluoromethyl)phenyl]-1H- isoIndole-1,3(2H)-dione	395.6
32	H ₂ N H ₃ C. O O F	3-bromo-4-{[(2- fluorophenyl)methyl]oxy]-5- (methyloxy)benzaldehyde thiosemicarbazone	413.3

	Q"		419.9
33	o to	2-[4-(3-chlorophenyl)piperazin-1- yl]-5-nitrobenzaldehyde thiosemicarbazone	419.9
34	G ON TOWN HE NH2	4-[[2-(3- chlorophenyl)ethyl]amino}-3- nitrobenzaldehyde thiosemicarbazone	378.9
35	H ₃ C CH ₃ N N N N N N N N N N N N N N N N N N N	(1E)-6,9-dimethyl-2,3,4,9- tetrahydro-1H-carbazol-1-one thiosemicarbazone	287.4
36	H ₂ N S HN. N	(2E)-1,1'-bi(cyclohexan)-1-en-2- one thiosemicarbazone	252.4
37	CONTRACTOR NAME OF THE PARTY OF	4-[[2-(4- chlorophenyl)ethyl]amino}-3- nitrobenzaldehyde thiosemicarbazone	378.9
38	H,C , , , , , , , , , , , , , , , , , ,	4-(diethylamino)-2-{[(4- fluorophenyl)methyljoxy}benzald ehyde N-{2-piperidin-1- ylethyl)thiosemicarbazone	486.7
39	H ₃ C N-N O-CH ₃	3,4-bis(methyloxy)benzaldehyde (1,1-dioxido-1,2-benzisothiazol- 3-y))(methy))hydrazone	360.4
40	H ₂ N NH HN N GI	(2E)-2-[(4-chlorophenyl)(5- chlorothien-2- yl)methylidene]hydrazinecarboxi midamide	314.2

41	,	2-(4-amino-2-oxo-1-propyl-1,2- dihydroquinolin-3-yl)-1H- benzimidazole-6-carbonitrile	344.4
42	NCO THE POST	4-amino-6-fluoro-7-({[4- (methyloxy)phenyl]methyl)amino)-3-[5-(4-methylpiperazin-1-yl)- 1H-benzimidazol-2-yl]quinolin- 2(1H)-one	528.6
43	CH,	6-chloro-3-(5-chloro-1H- benzimidazol-2-yl)-4-{[2- (dimethylamino)ethyl]amino}quin olin-2(1H)-one	417.3
44	H ₃ C N N N N N N N N N N N N N N N N N N N	4-amino-5-(1H-benzimidazol-2- yl)-1-methyl-1,7-dihydro-6H- pyrazolo(3,4-b]pyridin-6-one	281.3
45	O _{2N} CH ₃ O _{2N} CH ₃	5,5-dimethyl-4-methylldene-3- (2,4,6-trinitrophenyl)-1,3- oxazolidin-2-one	339.2
46	H ₃ C O CH ₃	5-methyl-2-[4- (methyloxy)phenyl]hexahydro- 1H-isoindole-1,3(2H)-dione	274.3
47	H ₉ C CH ₉	5-methyl-2-(4- methylphenyl)hexahydro-1H- isoindole-1,3(2H)-dione	258.3
48	H ₂ N N CHEH ₃	N-2(4-chlorophenyl)-6,6- dimethyl-1,6-dihydro-1,3,5- triazine-2,4-diamine	252.7

49		(7Z)-7-(turan-2-ylmethylidene)-3- phenyl-3,4-dihydro-2H- [1,3]thiazolo[3,2-a][1,3,5]triazin- (6(7H)-one	312.4
50	HO H H H	(3aR,9R,9aR)-6,7-dihydroxy-9- [3,4,5-tris(methyloxy)phenyl]- 3a,4,9,9a- tetrahydronaphtho[2,3-c]furan- 1(3H)-one	387.4
51	CI CH, CCH, CCH, CCH, CCH, CCH, CCH, CCH	G-chloro-2-(ethyloxy)-4-methyl-3- (4-nitrophenyl)-3a,4,9,9a- tetrahydro-3H-pyrrolo[2,3- b]quinoxaline	387.8
52	H _c C CH ₃	ethyl 2-(ethyloxy)-4-methyl- 3a,4,9,9a-tetrahydro-3H- pyrrolo[2,3-b]quinoxaline-3- carboxylate	304.4
53	H ₃ C H ₃ C H ₃ C H ₃ C	ethyl 4-([[2,5- bis(methyloxy)phenyl]amino)met hyl)-3,5-dimethyl-1H-pyrrole-2- carboxylate	333.4
54	SA IN FORM	1-{3-[(6-amino-5-nitropyridin-2- yl)amino]propyl)-4-{2- chlorophenyl-N-[(2S)-2- hydroxypropyl]-1H-pyrrole-3- carboxamide	473.9
55	C CH,	(4-methylphenyl)(5-nitro-2- piperidin-1-ylphenyl)methanone	325.4
56		(2S,5R)-N~1~-(4-methylphenyl)- 5-phenyl-N~2~-(2-pyridin-2- ylethyl)pyrrolidine-1,2- dicarboxamide	429.5

57	LO LA	2-[(3S)-3-(acetylamino)-2- oxopyrrolidin-1-yl]-N-[2-(4- fluorophenyl)ethyl]acetamide	322.4
58	COLINIAL COL	N-[2-(2,4-dichlorophenyl)ethyl]- 4-{{Z}-[(4,4- difluorocyclohexyl)imino][(3S)-3- methylpiperazin-1- yl]methyl]amino)benzamide	553.5
59	O-CH ₃	4-[4-(methyloxy)phenyl]-5- phenylisoxazole	252.3
60	FY THO HISTORY	methyl 4-{[4-(1-methylethyl)-2,3-dioxo-7-(trifluoromethyl)-3,4-dihydroquinoxalin-1(2H)-yl]methyl]benzoate	421.4
61	HO OH OH OH	(3beta,16beta)-3,14,16- trihydroxybufa-20,22-dienolide	403.5
62		2-(aminomethyl)-1-(2-pyridin-2- ylethyl)quinazolin-4(1H)-one	281.3
63	, E+E	ethyl 4-{[5-{3,4- bis(methyloxy)phenyl]-7- (trifluoromethyl)pyrazolo[1,5- a]pyrimidin-3- yl]carbonyl)piperazine-1- carboxylate	508.5
64	H _S C O O N O N O F F	5-[3,4-bis(methyloxy)phenyl]-3- (piperidin-1-ylcarbonyl)-7- (trifluoromethyl)pyrazolo[1,5- a]pyrimidine	435.4

65	H _G CO COLL	5-[3,4-bis(methyloxy)phenyl]-N- methyl-N-(2-pyridin-2-ylethyl)-7- (trifluoromethyl)pyrazolo[1,5- a]pyrimidine-2-carboxamide	486.5
66	H ₃ C N S OH	5-propyl-2-thien-2- ylpyrazolo[1,5-a]pyrimidin-7-ol	260.3

Table 35

Compound #	Structure	Source	Literature Reference	Patent Number
-	∠CH ₃ .H ₂ O			
-	Na S N CH ₃ .H ₂ O			
	Na S N CH ₃ .H ₂ O		 Lang, JM.; Touraine, JL.; Trepo, C. et al. 	
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1	OH	Aventis Pasteur	702-5.	
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288	O ₂ N O Na*	Aston University	Kinchington, D. et al. 4th Conf Retrovinuses Opportunistic infect (Jan 22-26, Washington DC) 1997, Abst .	

Table 35 Continued

Compound #	Structure	Source	Literature Reference	Patent Number
	CH ₃ CH ₃ O OH CH ₃		1) Fiedler-Nagy, C. et al.	
289		Roche	Agent Action 1989, 27(3- 4): 313-5.	EP 169571
290	OH CH3		1) Forest Laboratories,	
200		NFCR	Inc. Annual Report 1994.	WO 9517890
291	NH ₂	Sumitomo Pharmaceuticais		EP 248399
292	CH ₃ CH ₃ CH ₃	Aventis Pherma		
	o≼ ^{OH}	CALCUMA LIBRIDIA		EP 248734
293	N N N N N N N N N N N N N N N N N N N	May & Baker		EP 252682
294		Sumitomo Pharmaceuticals		EP 248399
295	F CH. NH2	Sumitomo Pharmaceuticals		EP 248399
296	O CH ₃ NH ₂	Sumitomo Pharmaceuticais		EP 248399
297	OF CH ₃ NH ₂	Sumitomo Pharmaceuticals		EP 248399
298	HO.N.CH3	Sumitomo Pharmaceuticals		EP 248399

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300 CH ₃ CH ₃ Sumitomo Pharmacouticais EP 248	
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302 CH ₃ CH ₃ Aveniis Pharma EP 2487	
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CH ₃ CH ₃ CH ₃ EP 24873	
Aventis Pharma EP 24973	4
307 NO. Aventis Pharma EP 24873	4
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309 H ₃ C CH ₃ Harbor Branch Found. US 475652	29

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373  Leo  EP 460032  EP 460032  AU 9057875  AU 9057875	3/2		Leo		EP 460032
373  Leo  EP 460032  Kricek, F. et al. Immunopharmacology 1907, 30(1). Let al. 31th and 100 an		ALC ALCOHOLOGICAL		1	
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991	HE THE SHEET ON LINE,	Beaufour-Ipsen	1) Carde, P. et al. Proc Amer Soc Clin Oncol 1991, 10: Abst 324.	AU 8810261
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992	о сн,	Roche		EP 510473
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996		Kyowa Hakko		EP 526840
997	HN-N CH ₃	Kyowa Hakko		EP 526840
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999	H ₃ C CN CH ₃	Aventis Pharma	-	EP 538783
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1000		Hayashibara	July 2, Stockholm) 1993, Abst 516.	EP 539196

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1196	3	Cell Therapeutics		WO 0268421

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1197	ĊH ₃	Pharmaceuticals		EP 248399
1198	CH ₃ CH ₃ CH ₃	Aventis Pharma		EP 248734
1199	F CH ₃ CH ₃	Sumitomo Pharmaceuticals		EP 248399
1200	F CH ₃	Sumitomo Pharmaceuticals		EP 248399
1201	OH ₂ NH ₂	Sumitomo Pharmaceuticals		EP 248399
1202	CH ₃	Sumitomo Pharmaceuticals		EP 248399
1203	Ho-N CH ₃	Sumitomo Pharmaceuticals		EP 248 399
1204	HO'N CH ₃ NH ₂	Sumitomo Pharmaceuticals		EP 248399
1205	O-N N-CH ₃	Sumitomo Pharmaceutloals		:P 248399
1206	EH ₃ NH ₂	Sumitomo Pharmaceuticals		P 248399

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1430	HO COM	Leo		WO 9737972
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1431	HO**CH	Leo		WO 9737972
1432	NG COL	Leo		WO 9737972
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BRIEF DESCRIPTION OF SEQUENCE LISTING

Description
Draft genome assembly from The Genome Science Center in British Colombia,
Canada of sequence from TOR2 isolate. TOR2_draft_genome_assembly_120403 Release 1
CDC SARS-CoV strain sequence. Entire nucleotide sequence (Urbani strain)
Group-specific coronavirus gene products
> Feline infectious peritonitis virus (FIPV)
3/4 = ORF 3b; $5/6 = ORF 3X$; $7/8 = ORF 3A$
> Canine coronavirus
9/10 = ORF 7b; 11/12 = ORF 7a
> Avian infectious bronchitis virus
13/14 = ORF 5b; 15/16 = ORF 5a; 17/18 = ORF 3a; 19/20 = ORF 3b
500 primers for left part
500 primers for right part
Forward primers from Table 4
Reverse primers from Table 4
Figure 9 primers
Figure 11 primers
Five primers from http://content.nejm.org/cgi/reprint/NEJMoa030781v2.pdf
PEP1 to PEP13
Extended PEP13
229E human coronavirus sequences
TGV sequences
PEDV sequences
Bovine coronavirus sequences
Murine hepatitis virus sequences
AIBV sequences
Primer sequences (forward)
Primer sequences (reverse)
Primer sequences (forward)
Primer sequences (reverse)
Primer sequences (forward)
Primer sequences (reverse)
Primer sequences (forward) F1-F48
Primer sequences (reverse) R1-R47
Primer sequences
Primer sequences
The nsp2 proteinase (3CL-PRO) sequence in SARS coronavirus
The nsp2 proteinases (3CLp) of avian IBV, MHV, and BCoV
Consensus nsp2 proteinases sequence
IG sequences from Figure 18
Expression construct of nSh in pCMVIII
Expression construct of nS in pCMVIII
Expression construct of nSh ATC in pCMVIII
Expression construct of nS ATC in pCMVIII
Expression construct of nS1h in pCMVIII
Expression construct of nS1 in pCMVIII Primers for cDNA amplification
Primers for CDNA amplification Primers for RT-PCR
Component sequences of Figure 23 (24 amino acids)
Component sequences of Figure 24 (≥4 amino acids) N-glycosylation sites within SEO ID NO: 6039
Liv-grycosyration sites within SEQ ID NO: 6039
Component conveness of Figure 25
Component sequences of Figure 25
Fragment of SEQ ID NO: 7188

7194	Amino acids 901-1005 of SEQ ID NO: 6042
7195	Amino acids 1144-1201 of SEO ID NO: 6042
7196	Amino acids 1144-1196 of SEQ ID NO: 6042
7197-7199	Membrane fusion peptide regions
7200-7206	NadA-based polypeptides
7207-7223	N-glycosylation sites within SEQ ID NO: 6042
7224-7231	Slippage region
7232	Orf lab polyprotein
7233-7244	Orflab polyproteins
7245-7247	Offiab polyproteins
7248-7253	X ₂ sequences for SEQ ID NOS 7233-7244
7254	Orf1ab polyproteins
	Zinc binding region 2 site
7255-7271	N-glycosylation sites in SEQ ID NOS: 6040-41,6043,6045-46,6050-51
7272-7291	Polypeptides and polynucleotides
7292-7293	Intergenic sequences
7294-7301	Nucleotides from 5' end of SARSV genome followed by intergenic sequence
7302-7306	
7307-7308	Fragments of SEQ ID NO: 6042
7309	NadA sequence
7310-7311	NadA leader sequences
7312-7315	Amino acid sequencess from NadA
7316-7324	PCR primers
7325-7330	Primers
7331	CCACC sequence
7332-7336	3' UTR forward primers
7337-7341	3' UTR reverse primers
7342-7352	3' UTR probes
7353-7362	5' UTR forward primers
7363-7373	5' UTR reverse primers
7374-7385	5' UTR probes
7386	
7387	Conserved octanucleotide
7388	Reverse complement of SEQ ID NO: 7293
7389	Intergenic sequence
7390	Poly T
	Stem-loop sequence
7391-7392	Poly-glycine linkers
7393	Poly-histidine tag
7394	Nucleocapsid epitope site
7395	Antisense primer
7396-7397	Probes
7398-7399	Antigenic fragments of SEQ ID NO: 6042
7400-7639	1-epitope analysis of SEO ID NO: 6039
7640-7800	T-epitope analysis of SEO ID NO: 6040
7801-8040	T-epitope analysis of SEQ ID NO: 6041
8041-8280	T-epitope analysis of SEO ID NO: 6042
8281-8486	T-epitope analysis of SEO ID NO: 6043
8487-8665	T-epitope analysis of SEO ID NO: 6044
8666-8820	T-epitope analysis of SEO ID NO: 6045
8821-9018	T-epitope analysis of SEQ ID NO: 6046
9019-9131	T-epitope analysis of SEO ID NO: 6047
9132-9308	T-epitope analysis of SEQ ID NO: 6048
9309-9437	T-epitope analysis of SEQ ID NO: 6049
9438-9538	T-epitope analysis of SEQ ID NO: 6050
9539-9752	T-epitope analysis of SEQ ID NO: 6052
9753-9763	Primers for spike protein amplification
9764-9765	Primers for spike protein amplification, particularly fragments of spike N-glycosylation sites within SEQ ID NO: 6039
9766-9779	
	Cleavage products for ORF1ab (Table 10)

9780-9782	Forward primer, reverse primer, probe
9783-9784	Lysine-rich region
9785-9798	Oligonucleotides used for S.cerevisiae expression
9799-9802	Sequences from Figures 65 & 66
9803-9882	Primers for E.coli cloning
9883-9885	BCV nucleotide sequences for Figures 3A, 3B, 3C
9886-9891	BCV amino acid sequences for Figures 4A, 4B, 4C, 4D, 4E, 4F
9892	BCV 5' UTR
9893	BCV 3' UTR
9894-9896	MHV nucleotide sequences for Figures 3A, 3B, 3C
9897-9902	MHV amino acid sequences for Figures 4A, 4B, 4C, 4D, 4E, 4F
9903-9904	AIBV nucleotide sequences for Figures 3A, 3B
9905-9909	AIBV amino acid sequences for Figures 4A, 4B, 4D, 4E, 4F
9910	AIBV 5' UTR
9911	AIBV 3' UTR
9912-9913	HOBMPRO, HOBHEGA nucleotide sequences for Figures 3B, 3C
9914-9918	Human CoV amino acid sequences for Figures 4A, 4B, 4C, 4E, 4F
9919	HCoV-OC43 5' UTR
9920	HCoV-OC43 3' UTR
9921-9923	pCMVKm2 vectors
9924-9926	Codon-optimised N, M and E sequences
9927	BNI-1
9928-9959	Constituent amino acid sequences ≥4aa inferred from SEQ ID NO: 9927
9960	ORF1ab variant
9961	ORF1a variant
9962	Spike variant
9963	Membrane variant
9964	Nucleocapsid variant
9965-9966	Short ORFs
9967	FRA complete genome

CLAIMS

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1. An isolated polypeptide of the SARS virus.

- 2. The polypeptide of claim 1, wherein the polypeptide is a Spike (S) polypeptide, an Env (E) polypeptide, a Membrane (M) polypeptide, a hemagglutinin-esterase polypeptide (HE), a nucleocapsid (N) polypeptide, a ORF1a polypeptide, a ORF1ab polypeptide, a Proteolytic fragment of a ORF1ab polypeptide.
- 3. The polypeptide of claim 1, wherein the polypeptide comprises an amino acid sequence selected from the group consisting of SEQ ID NO⁵: 6039, 7232, 9766, 9767, 9768, 9769, 9770, 9771, 9772, 9773, 9774, 9775, 9776, 9777, 9778, 9779, 6042, 6043, 6044, 6045, 6046, 6047, 6048, 6049, 6050 or 6052.
- 4. The polypeptide of claim 1, wherein the polypeptide comprises an amino acid sequence having >75% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO^S: 6042, 6043, 6044, 6045, 6046, 6047, 6048, 6049, 6050, 6052, 9766, 9767, 9768, 9769, 9770, 9771, 9772, 9773, 9774, 9775, 9776, 9777, 9778, 9779, 9997, 9998, 10149, 10316, 10338, 10339, 10340, 10341, 10342, 10532, 10533, 10571, 10572, 10573, 10574, 10575, 10576, 10577, 10578, 10579, 11561, 11562, 11618, 11619, 11620, 11627, 11630, 11633 & 11636.
- 5. The polypeptide of claim 1, wherein the polypeptide comprises a fragment of at least 10 consecutive amino acids of an amino acid sequence selected from the group consisting of SEQ ID NO⁸: 6042, 6043, 6044, 6045, 6046, 6047, 6048, 6049, 6050, 6052, 9766, 9767, 9768, 9769, 9770, 9771, 9772, 9773, 9774, 9775, 9776, 9777, 9778, 9779, 9997, 9998, 10149, 10316, 10338, 10339, 10340, 10341, 10342, 10532, 10533, 10571, 10572, 10573, 10574, 10575, 10576, 10577, 10578, 10579, 11552, 11561, 11562, 11618, 11619, 11620, 11627, 11630, 11633 & 11636.
- A polypeptide comprising an amino acid sequence having >80% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO⁸: 6042, 6043, 6044, 6045, 6046, 6047, 6048, 6049, 6050, 6052, 9766, 9767, 9768, 9769, 9770, 9771, 9772, 9773, 9774, 9775, 9776, 9777, 9778, 9779, 9998, 10149, 10316, 10338, 10339, 10340, 10341, 10342, 10532, 10533, 10571, 10572, 10573, 10574, 10575, 10576, 10577, 10578, 10579, 11552, 11561, 11562, 11618, 11619, 11620, 11627, 11630, 11633 & 11636.
- 7. A polypeptide comprising an amino acid sequence that comprises a fragment of at least 10 consecutive amino acids of an amino acid sequence selected from the group consisting SEQ ID NO⁸: 6042, 6043, 6044, 6045, 6046, 6047, 6048, 6049, 6050, 6052, 9766, 9767, 9768, 9769, 9770, 9771, 9772, 9773, 9774, 9775, 9776, 9777, 9778, 9779, 9997, 9998, 10149, 10316, 10338, 10339, 10340, 10341, 10342, 10532, 10533, 10571, 10572, 10573, 10574, 10575, 10576, 10577, 10578, 10579, 11552, 11561, 11562, 11618, 11619, 11620, 11627, 11630, 11633 & 11636.

8. A polypeptide comprising an amino acid sequence having >80% sequence identity to SEQ ID NO: 6042, and/or comprising an amino acid sequence that comprises a fragment of at least 10 consecutive amino acids of SEQ ID NO: 6042, wherein the polypeptide is in the form of a trimer.

- 5 9. Nucleic acid encoding the polypeptide of any one of claims 1 to 8.
 - Nucleic acid according to claim 9, comprising a nucleotide sequence selected from the group consisting of SEQ ID NO⁸: 7191, 7273, 7275, 7277, 7279, 7281, 7283, 7285, 7287, 7289, 7291, 7293, 9968, 10066, 10084, 10299, 10505, 11323, 11563, 11639 & 11640.
 - 11. A polynucleotide comprising a nucleotide sequence having >80% sequence identity to the nucleic acid of claim 9 or claim 10.
 - 12. A polynucleotide comprising a fragment of at least 10 consecutive nucleotides of the nucleic acid of claim 9 or claim 10.
 - 13. Antibody that recognizes the polypeptide of any one of claim 1 to 8.
- The antibody of claim 13, wherein said antibody recognizes the polypeptide comprising
 the amino acid sequence of SEQ ID NO: 6042 or a fragment thereof.
 - 15. The antibody of claim 14, wherein said antibody recognizes the polypeptide comprising the amino acid sequence of SEQ ID NO: 6042 or a fragment thereof in trimeric form.
 - 16. The antibody of claim 13, wherein the antibody is a monoclonal antibody,
 - 17. The antibody of claim 13, wherein the antibody is a human antibody,
- 20 18. An immunoassay for detecting a SARS virus antigen in a sample, comprising the step of contacting the sample with the antibody of any one of claims 13 to 17.
 - 19. An immunoassay for detecting an antibody against a SARS virus antigen in a sample, comprising the step of contacting the sample with the polypeptide of any one of claims 1 to 8.
- 20. A method of detecting an antibody against a SARS virus antigen in a sample comprising contacting said sample with the polypeptide of any one of claims 1 to 8, under conditions suitable for binding said polypeptide to said antibody, if present, and detecting the binding of said polypeptide to said antibody.
 - 21. A method for detecting a SARS virus antigen in a sample comprising contacting said sample with the antibody of any one of claims 13 to 17, under conditions suitable for binding said antibody to said antigen, if present, and detecting the binding of said antibody to said antigen.

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22. A vaccine for the treatment or prevention of severe acute respiratory syndrome (SARS), comprising an inactivated SARS virus, a killed SARS virus, an attenuated SARS virus, a split SARS virus preparation, or at least one purified SARS virus antigens.

- 23. The vaccine of claim 22, comprising a purified polypeptide according to any one of claims 1 to 8.
- 24. The vaccine of claim 22 or claim 23, wherein the antigen is a purified SARS virus antigen in the form of a VLP.
- 25. The vaccine of any one of claims 22 to 24, further comprising an adjuvant.
- The vaccine of claim 25, wherein the adjuvant is an aluminium salt or is MF59.
- 10 27. The vaccine of any one of claims 22 to 26, comprising more than one SARS virus antigen.
 - 28. The vaccine of claim 27, wherein the antigens are selected from S, E, N and M.
 - 29. The vaccine of claim 22, comprising an inactivated SARS virus.
 - 30. The vaccine of claim 29, wherein said virus is inactivated by chemical or physical means.
- 15 31. The vaccine of claim 30, wherein said inactivation comprises treatment of the virus with an effective amount of one or more of the following agents selected from the group consisting of detergents, formaldehyde, formalin, β-propriolactone, and UV light.
 - 32. The vaccine of claim 30, wherein said inactivation comprises treatment of the virus with an effective amount of one or more of the following agents selected from the group consisting of methylene blue, psoralen and carboxyfullerene (C60).
 - 33. The vaccine of claim 30, wherein said inactivation comprises treatment of the virus with an effective amount of one or more of the following agents selected from the group consisting of binary ethylamine, acetyl ethyleneimine and gamma irradiation.
 - 34. The vaccine of claim 31, wherein said inactivation comprises treatment with β propriolactone.
 - 35. The vaccine of claim 34, wherein said β -propriolactone is used at a concentration of 0.01 to 0.5%.
 - 36. The vaccine of claim 34, wherein said β -propriolactone is used at a concentration of 0.5 to 0.2%.
- 30 37. The vaccine of claim 34, wherein said β-propriolactone is used at a concentration of 0.025 to 0.1%.

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38. A method of inactivating SARS virus comprising exposing the virus to an inactivation agent for 12 to 24 hours at refrigeration temperatures followed hydrolysis of any residual inactivating agent by elevating the temperature for three hours.

- 39. The method of claim 38, wherein the inactivation agent is β-propriolactone.
- 5 40. The method of claim 38, wherein the refrigeration temperature is between 0°C and 8°C.
 - 41. The method of claim 38, wherein the elevated temperature is between 33°C and 41°C.
 - 42. A method for making an inactivated SARS vaccine comprising:
 - a. innoculating a mammalian cell culture with SARS virus;
 - b. cultivating the infected cells;
 - c. harvesting SARS virus containing supernatant:
 - d. inactivating the SARS virus; and
 - e. purifying the inactivated SARS virus.
 - 43. The method of claim 42, wherein said mammalian cell culture is derived from one or more of the cell types selected from the group consisting of fibroblast cells, endothelial cells, hepatocytes, keratinocytes, immune cells, mammary cells, smooth muscle cells, melanocyte cells, neural cells, prostate cells, renal cells, skeletal cells, liver cells, retinoblast cells and stromal cells.
 - 44. The method of claim 42, wherein said mammalian cell culture is derived from a cell culture selected from the group consisting of human cells, non-human primate cells, HeLa cells, human diploid cells, fetal rhesus lung cells, human embryonic kidney cells, VERO cells, horse cells, cow cells, sheep cells, dog cells, cat cells or rodent cells.
 - 45. The method of claim 42, wherein said mammalian cell culture is derived from VERO cells or fetal rhesus kidney cells.
 - 46. The method of claim 42, wherein said mammalian cells are cultured in serum free media.
- 25 47. The method of claim 42, wherein said mammalian cells are cultured in protein free media.
 - 48. The method of claim 42, wherein said inoculating step comprising absorbing the SARS virus onto the cell culture for 60 to 300 minutes.
 - The method of claim 42, wherein said inoculating step is conducted at 25°C to 40°C.
- 50. The method of claim 42, wherein said purification step comprises one or more of the treatments selected from the group consisting of gradient centrifugation, ultracentrifugation, continuous-flow ultracentrifugation, chromatography, polyethylene glycol precipitation, and ammonium sulfate precipitation.

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51. The method of claim 42, wherein said purification step comprises one or more of the treatments selected from the group consisting of ultrafiltration and dialfiltration.

- 52. The method of claim 50, wherein said chromatography treatment includes one or more of the chromatography treatments selected from the group consisting of ion exchange
- 5 chromatography, size exclusion chromatography, and liquid affinity chromatography.
 - 53. The method of claim 52, wherein said chromatography treatment includes use of one more chromatographic resins selected from the group consisting of an an anionic resin and a cationic resin.
- 54. The method of claim 52, wherein the ion exchange chromatography treatment includes a first step using a strong anion exchange resin and a second step using a strong cation exchange resin.
 - 55. The method of claim 50, wherein said gradient centrifugation purification step comprises density gradient centrifugation.
 - 56. The method of claim 42, wherein said purification step comprises a first step of chromatography purification and a second step of gradient centrifugation.
 - 57. The method of claim 56, wherein said first chromatography purification step comprises liquid affinity chromatography.
 - The method of claim 56, wherein said second gradient centrifugation step comprises density gradient centrifugation.
- A single-stranded oligonucleotide comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS: 21-6020, 6076-6568, 6586-6587, 7292-7301, 7325-7328, 7332-7352, 7353-7385, 10235-10298, 10352-10504, 10580-11322 and 11325-11551.
 - A single-stranded oligonucleotide comprising the complement of the oligonucleotide of claim 59.
- 25 61. The oligonucleotide of claim 59 or claim 60, comprising 10-30 nucleotides.
 - 62. The oligonucleotide of claim 61, comprising the nucleotide sequence of SEQ ID NO: 7292, SEQ ID NO: 7293, the complement of SEQ ID NO: 7292 or the complement of SEQ ID NO: 7293.
- 63. A kit comprising primers for amplifying a template sequence contained within a SARS virus nucleic acid target, the kit comprising a first primer and a second primer, wherein the first primer comprises a sequence substantially complementary to a portion of said template sequence and the second primer comprises a sequence substantially complementary to a portion of the

complement of said template sequence, wherein the sequences within said primers which have substantial complementarity define the termini of the template sequence to be amplified.

- 64. The kit of claim 63, wherein the template sequence is contained within SEQ ID NO: 1 and/or SEQ ID NO: 2.
- 5 65. The kit of claim 63 or claim 64, wherein the first primer comprises a fragment of 8 or more nucleotides of SEQ ID NO: 1, and the second primer comprises a fragment of 8 or more nucleotides of the complement of SEQ ID NO: 1.
 - 66. The kit of claim 63 or claim 64, wherein the first primer comprises a fragment of 8 or more nucleotides of SEQ ID NO: 2, and the second primer comprises a fragment of 8 or more nucleotides of the complement of SEQ ID NO: 2.
 - 67. The kit of claim 63, wherein the first primer is an oligonucleotide according to any one of claims 59 to 62 and the second primer is an oligonucleotide according to any of claims 59 to 62.
 - 68. The kit of any one of claims 63 to 67, further comprising a labeled probe that comprises either a fragment of 8 or more nucleotides of SEQ ID NO: 1 and/or SEQ ID NO: 2, or the complement of said fragment, which fragment is located within the template sequence.
 - 69. The kit of any one of claims 63 to 68, wherein the first primer and/or the second primer comprises a nucleotide sequence selected from the group consisting of SEQ ID NOS: 21-6020, 6076-6568, 6586-6587, 7292-7301, 7325-7328, 7332-7352, 7353-7385, 10235-10298, 10352-10504, 10580-11322 and 11325-11551.
- 70. The kit of any one of claims 63 to 68, wherein the first primer and/or the second primer comprises the complement of a nucleotide sequence selected from the group consisting of SEQ ID NOS: 21-6020, 6076-6568, 6586-6587, 7292-7301, 7325-7328, 7332-7352, 7353-7385, 10235-10298, 10352-10504, 10580-11322 and 11325-11551.
 - 71. A method of detecting the presence of SARS virus in a sample comprising providing a sample suspected of containing a SARS virus nucleic acid target, amplifying a template sequence contained within said SARS virus nucleic acid target with the kit of any one of claims 63 to 70, and detecting the amplified template sequence, wherein the presence of the amplified template sequence indicates the presence of SARS virus in said sample.
 - 72. The method of claim 71, wherein said amplifying is accomplished using polymerase chain reaction, transcription mediated amplification, reverse transcription PCR, ligase chain reaction, strand displacement amplification or nucleic acid sequence-based amplification.
 - 73. A double-stranded RNA molecule with a length from about 10 to about 30 nucleotides which is able to inactivate the SARS coronavirus in a mammalian cell.

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74. The double-stranded RNA of claim 73, wherein the sequence of one of the strands is at least 90% identical to a target sequence, wherein the target sequence is a fragment of SEQ ID NO: 1 and/or SEO ID NO: 2.

- 75. The double-stranded RNA of claim 73 or claim 74, wherein the target sequence comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS: 7292, 7293, 7294, 7295, 7296, 7297, 7298, 7299, 7300 and 7301.
- 76. The double-stranded RNA of any one of claims 73 to 75, comprising at least one modified nucleotide.
- 77. A method for treating a patient suffering from SARS, comprising: administering to the patient a therapeutically effective dose of a molecule of less than 1000 g/mol.
- 78. The method of claim 77, wherein the molecule has an aromatic region and greater than one heteroatom selected from O, S, or N.
- 79. A method for treating a patient suffering from SARS, comprising: administering to the patient a therapeutically effective dose of a compound selected from: a nucleoside analog, a peptoid, an oligopeptide, a polypeptide a protease inhibitor, a 3C-like protease inhibitor, a papain-like protease inhibitor, or an inhibitor of an RNA dependent RNA polymerase.
- 80. A method for treating a patient suffering from SARS, comprising: administering to the patient a steroidal anti-inflammatory drug in combination with at least one antiviral compound.
- 81. A method for treating a patient suffering from SARS, comprising: administering to the patient a therapeutically effective dose of a compound selected from: acyclovir, gancyclovir, vidarabidine, foscamet, cidofovir, amantidine, ribavirin, trifluorothymidine, zidovudine, didanosine, zalcitabine, an antiviral compound listed in Table 1; an antiviral compound listed in Table 2; or an interferon.
 - 82. The method of claim 81, wherein the interferon is an interferon-α or an interferon-β.
- 25 83. The method of any one of claims 77 to 82, wherein the molecule or compound is delivered by inhalation.
 - 84. A method of identifying a therapeutically active agent comprising the steps of: (a) contacting a therapeutically active agent with a cell infected with the SARS virus; (b) measuring attenuation of a SARS related enzyme.
- 30 85. A viral vector or particle for in vivo delivery of a nucleic acid of claim 9 or claim 10.
 - 86. The viral vector of claim 85, wherein the vector is an adenovirus vector, a poxvirus vector or an alphavirus vector.
 - 87. An alphavirus replicon particle comprising one or more SARS viral antigens.

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88. The replicon particle of claim 87, wherein said SARS viral antigen is a spike protein.

- 89. The replicon particle of claim 87, wherein said particle comprises a replicon derived from Venezuelan Equine Encephalitis (VEE) and further comprises an envelope derived from Sindbus virus (SIN) or Semliki Forest Virus (SFV).
- 5 90. A vaccine comprising one or more SARS virus antigens and one or more respiratory virus antigens.
 - 91. The vaccine of claim 90, wherein said respiratory virus antigens are selected from the group consisting of influenza virus, human rhinovirus (HRV), parainfluenza virus (PIV), respiratory syncytial virus (RSV), adenovirus, metapneumovirus, and rhinovirus.
- 10 92. The vaccine of claim 91, wherein said respiratory virus antigen is from influenza virus.
 - 93. The vaccine of claim 90, wherein said respiratory virus antigen is from a coronavirus other than the SARS virus.
 - A polypeptide comprising an immunogenic, surface exposed fragment of the amino acid sequence SEQ ID NO: 6042.
- 15 95. The polypeptide of claim 94, wherein said fragment does not include the last 50 amino acids of the C-terminus of SEQ ID NO: 6042.
 - 96. The polypeptide of claim 94, wherein said fragment does not include a transdomain region of SEQ ID NO: 6042.
 - 97. The polypeptide of claim 94, wherein said fragment does not include a C-terminus cytoplasmic domain of SEQ ID NO: 6042.
 - 98. The polypeptide of claim 94, wherein said fragment does not include a N-terminus signal sequence.
 - 99. An isolated polynucleotide comprising a nucleic acid sequence selected from the group consisting of SEQ ID NOS: 9968 and 10066.
- 25 100. The polynucleotide of claim 99, wherein the polynucleotide comprising a nucleic acid sequence having > 80% sequence identity to a polynucleotide sequence selected from the group consisting of SEQ ID NOS: 9968 and 10066.
 - 101. An isolated polynucleotide comprising a fragment of at least 15 consecutive nucleic acids of a nucleic acid sequence selected from the group consisting of SEQ ID NOS: 9968 and 10066 and wherein said fragment does not consist entirely of SEQ ID NO: 10033.
 - 102. An isolated polypeptide comprising an amino acid sequence encoded by any one of claims 99 101.

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103. The polypeptide of claim 102, comprising an amino acid sequence selected from the group consisting of SEQ ID NOS: 9969 – 10032, 10067, and 10015.

- 104. The polypeptide of claim 103, wherein the amino acid sequence is selected from the group consisting of SEQ ID NOS: 9997, 9998 and 10015.
- 5 105. An expression construct for recombinant expression of a SARS virus spike protein wherein said construct comprises a nucleic acid sequence selected from the group consisting of SEQ ID NOS: 6578 – 6583.
 - 106. A mammalian cell line stably expressing a SARS viral antigen.
 - 107. The cell line of claim 106, wherein said cell line is a Chinese Hamster Ovary (CHO) cell.
- 10 108. The cell line of claim 106, wherein the SARS viral antigen is a spike protein or fragment thereof.
 - 109. The cell line of claim 106, wherein the spike protein is truncated to remove the transmembrane sequence.
- 110. A method of identifying a therapeutically active agent comprising the steps of: (a) contacting a therapeutically active agent with a buffer comprising SARS enzyme; and (b) measuring attenuation of the SARS enzyme.
 - 111. The method of claim 110 wherein the SARS enzyme is a SARS protease.
 - 112. The method of claim 111 wherein the buffer further comprises a peptide with a SARS protease cleave site.
- 20 113. The method of claim 110 wherein the measurement is made by the measurement of fluorescence.
 - 114. A vaccine of one of claims 22 to 37, and 90 to 93 further comprising an adjuvant.
 - 115. The vaccine of claim 114 wherein the adjuvant is a SMIP.
- 116. The vaccine of claim 115 wherein the SMIP compound is selected from the group consisting of an acylpiperazine, a tryptanthrin, an indoledione, a tetrahydroisoquinoline, a benzocyclodione, an amino azavinyl compound, a thiosemicarbazone, a lactam, an aminobenzimidazole quinolinone, a hydropthalamide, a benzophenone, an isoxazole, a sterol, a quinazolinone, a pyrole, an anthraquinone, a quinoxaline, a triazine, an benzazole, and a pyrazolopyrimidine, or a pharmaceutically acceptable salt, ester, or prodrug thereof.
- 30 117. A method of vaccinating a subject comprising administering a vaccine of one of claims 22 to 37, and 90 to 93.
 - 118. The method of claim 117 further comprising administering a SMIP.

119. A method for treating a patient of one of claims 77 to 82 further comprising administering at least one SMIP compound.

120. A method for treating a patient of one of claims 77 to 82 further comprising administering at least one SMIS compound.

FIGURE 1

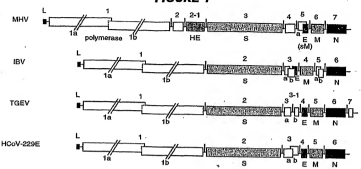


FIGURE 2

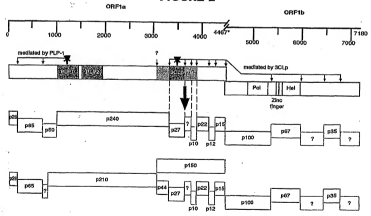


FIGURE 3

FIGURE 3A

												s	ection 1
	(1)			10			20			,30			4
BCV N	(1)	ATG	TCTT	TTA	CTC	TGG	T-AA	GCA	AT	CCA	GTA	GTAC	ÄGCG
MHV N	(1)	ATG	TCTT	TTG	TTCC	TGG	GCAA	GAA	AATC	CCG	GTG	GCAC	AAGC
Avian infectious bronchitis N	(1)												
Consensus	(1)	ATG	TCTT	TT		A ection 2							
	(44)	44	.50	0		6	0		.70				8
BCV N	(41)	cer	TOGG	AAA	rčæ	TCT	GGTA	ATG	CCAT	COL	TAA	AJ	
MHV N	(44)	COT	CTGT	AAA	cecie	Gor	CCTA	N'TG	ר מ מים	COL	CAA	a a c	ACCA
Avian infectious bronchitis N	(1)							Maria.		40.00	~300	2000	
Consensus	(44)	CCT	TG	AAA	CG	CT	GGTA	ATG	GAT	CCT	ממי	G	
									- 11.		****		ection 3
	(87)	87			4	00		4.	10				12
BCV N			aaac	Car			Acton	300	സ്ക്രീര്	in a m	Cmin	34 h z	CCAG
MHV N	(87)	TITICS	ACOT	CAC	330	000	Arion	TO C	NOO	77.7	224		ATAG
Avian infectious bronchitis N	(1)	14.00	SAME.	2.57 C	- APT (0.00	7000	CHA	X 3. 24		MAG		TCAG
Consensus	(87)		GGC				AGCG						
	(01)		GGC	GA (CA	CCG	MGCG	T	MAGE	MAT	1		CAG ection 4
	(130)	120			,140 ,150 ,160								
BCV N	(100)	100	Contract.			1- /250	110	OU TOTAL	USE CON	1	60	SPERM	17
BCV N MHV N	(110)	34.	2200	CAG	31.7	ACU	CAAG	UAA	ACTO	CTA	CTT	CTCA	GCTA
Avian infectious bronchitis N	(130)	2.3	MOAA	GGA	A 1.07	VG CC	AAPI	CAG	ACTO	CAA	CTA	CŢÇ	硬
	(430)	CTE	2000	GC G	CA	11 U-	AT	4.0	A C L F	GGA	GGA	CCA	AA
Consensus	(130)	GGC	AGAA	G G	CTC	4 66	AAG	CAA	ACT	e A	CTA		
	(173)	477		180			190					S	ection 5
BCV N					aleta E	P2-22-2		A 20 m A		00	50 F. 16	*: 17*21F.17	21
MHV N	(101)	CAT	CAGG	AGG	AA.	STT	GTAC	CGT	ACIT	T - T	CTŢ	geri	CTCT
Avian infectious bronchitis N	(71)	UAA	CTCC	- 666	JAG.	GUG	GTTC	ecc	ATU	E - I	CCT	ggra	Tron
Consensus	(71)	CAC	CAAA	- G@	TAGC	a G', Ti C	AUCT	GGA	AATEC	CAT	CIT	OOTA	TCAA
Consensus	(173)	CA	CA	GGG	GAG'	GT	GT C	CC	A TA	C T	CTT		TTCT
	****											8	ection 6
	(216)		290			230			240 ·				25
BCV N	(203)	GAA	TTA-	C	rca (TTT	CAAA	AAG	GAAZ	GĠA	CTT	TGAZ	TTTG
MHV N	(212)	GCA	TTA-	C	CCAC	TTC	CAAA	AGG	GAAZ	GGA	GTT	TCAC	TTTG
Avian infectious bronchitis N	(113)	CCA	TAAA	GGC	DAAC	AAA	CTAA	ATC	CACC	TGC	ACC	TAAC	TTTG
Consensus	(216)	GCA											
				s	ection 7								
	(259)	259			270			280			290		30
BCV N	(243)	AGA	GĠĠĀ	CAA	SGT(TOC	CTAT	TGC	ACC	GGA	GEC	CCAC	CTAC
MUAN M	(Z5Z)	AGA	AGGA	CAA	3GA(ST GC	CTAT	$\mathbf{T}\mathbf{G}\mathbf{C}$	CAAT	CCA	ATC	CCC	ب سينيان
Avian infectious bronchitis N	(155)	A	ACGT	AGT	GGT	TTC	CTGA	TAA	TGAZ	AAT	CTT	AAA	ATAG
Consensus	(259)	AGA	AGGA	CAAC	CTC	TOO	CTAT	TGC	7.7	GGA	T.C	C C D C	Canya

		(302)	303			310									- Seci	ion 8
	BCV N	(286)	0.02	aan	2000	110	25 3 5	20.00	320	*********		3	30			34
	BCV N	(295)	d s	17.25	200	1000	7. FL	TGC	TA	AGA	ÇA	CAA	CAG	ACG	TTCT	TTT.
Avian infectious	MHV N s bronchitis N	(106)	Oxo	CAL	THE REAL	AUUA	LAI	rGG	TA.	raga	OΑ	CAA	CCG	CCG	TTCT	PTT
	Consensus	(002)	GAG	CA	AAC	3GGG	TAC	TGO	TA	AGA	CA	CAA	CAG	CG	TTCT	PTT.
															- Sect	ion 9
	20111	(345)	345	3	50	*******		360			37	ro				38
	BCV N MHV N	(329)	AAA	CAG	CCC	ATG	GCA	- A C	CAC	CGT	CA	ACT	CCT	CCC	ACCA	
Avian Infantion	MHV N bronchitis N	(338)	AAA	CAC	cro	ATG	GGC	- AC	CAC	AAG	CA	ATT	ACT	GCC	CALLA	da
Avian Infectious																
	Consensus	(345)	AAA	CAG	CTC	ATG	GC	À	CAC	AA	CA	ATT	CT	GCC	CGA	יממי
	BCV N MHV N	(388)	388				400			410)			420		42
	BCV N	(371)	ATT	TATT	ACh	ATC	TTG	CAA	CAC	GÁĆ	"COV	D.T.	WAC.	7.8.5	do dos	200
Andrea torresta	MHV N bronchitis N	(380)	ATT	TTT	Аdд	ATC	TIG	GCA	CAC	GGG	CC	AT	700	200	CONC	5
Avian infectious	bronchitis N	(275)	ATT	TCT	ATT	ACA	ĈТĠ	GAA	CAC	GAIC	CAC	CC	10.1			
	Consensus	(388)	ATT	TTT	ACT	ATC	TTG	GAA	CAG	GAC	C (יייי	307	CAN	CIGN	PF
													JC 1	GMM	Sectio	- 44
		(431)	431			440			.450	` _			160			
	BCV N MHV N bronchitis N	(414)	TGG	CAC	CGA	SEA 13	raa.	cala	7 6 7	12144	rim e	150	46U	2737175		47
	MHV N	(423)	TCC	AGA	CAG	CAT	100	T CO	W (2.0	244	4.5	100	CG	CLA	JTAAC	CAC
Avian infectious	bronchitis N	(318)	GCC	TGA	TTC	TCA	A rain	min in	77 70 70	200	C 11	GG.	11.0	JAA	ACAGC	CAI
	Consensus	(431)	TGG	GA	c	TATE	rca.	100	ALC: U	CER	G T C	199	TG	garge (CTAAC	GG:
									161	CII	CIG	-GG;	rre	CTA	TAAC	CA
		(474)	474		480			490							Section	12
	BCV N	(457)	achie	dzin	CM'C	Or a my	242	490	- 35.4	Carlo Fee V	5	00	Pentille			516
	MHV N	(466)	GOO!	CAL	Mary C		ALC:	CCG	GCI	GAC	АТТ	CIL	'CA'	rcg	GACC	CAF
Avian infectious	MHV N bronchitis N	(361)	COR	C N m	ACU Ama	44.7	LUC	egc	TCT	GAT.	ATI	GE	GA.	AÀG	GACC	CAN
	Consensus	(474)	GCI	GAT	GTC	AATA	rcc	CG	TCT	GAC	TTA	GTC	GA?	CAGO	GACC	CAA
		(517)	F42												Section	13
	BCV N MHV N	(517)	01/ ©300	-	95230	a record	530	and the		54	0					559
	MEN N	(500)	GTAC	3CG	AIG	AGGC	TA	TTC	CGA	CTA	- GG	בידק	COC	cci	GGCA	cde
Avian infectious	MHV N bronchitis N	(404)	CCAC	ATC:	ATC	agge	TA	PTC	CTA	CTA	- GG	TTI	GCC	occ	GGCA	CGG
Avian infectious	Concensus	(404)	ATA	AGT	TŢĢ,	AT-C	AA.	(AC	CCA	CTA	CGG	TTI	TCI	GAJ	GG	
	Consensus	(317)	GT'AC	3 2	AT'G	AGGC	TA	rrc	C A	CTA	GG	TTT	. Co	rooi	GGCA	CGG
															Section	14
		(560)	560			570			,58	30			590			
	BCV N MHV N	(542)	TACI	CCC	CTC	AGGC	TT	CT	ATA	TTG	ÀG	CCT	-	CAN	Camia	TOP
hidaa lufa st	MHV N bronchitis N	(551)	TATI	GC	TTC	AGGG	CT)	rrr)	ATG	TTG	AG	COT	d'm'c	CA A	COMO	100
Avian infectious																
	Consensus	(560)	TA T	CC	CTC	AGGG	TT	TT	. Tr	TTCI	ם מ	a car		22.0	CORC	110

											Sec	lion 15
		(603)	603	,610)	6	20		,63	0		64
	BC/ N	(585)	TCC	PAATTO	CAGA	TCTAC	TTCA	CCC	GCA-	TG	CAGTA	AGC
Avian infectious												
	Consensus	(603)	ACC:	r CT	CAGA	TCTGG	TTCG	CG.	GCA	ATC	AGTG	SAGO
											Sect	ion 16
		(646)	646			660		67	'n			
	BCV N	16251	mam	AGTGCA	OC 2T	Clayba	A CTY		1907-7-1	min dem	187111 - I	68
Avian infectious	MHV N	(636)	AAA		- 2	acac.	- Cma	07.5	TOTAL	THET	GCAAC	CAGA
Avian infectious	bronchitis N	(514)	PCIA:	CARCO	COM	GTAC-	- Case	CAA	ale MC	THEC	ACCAC	SCGC
	Consensus'	(646)	TCA	r ac	G AM	GLAG-	ILA	GWA.	PACC	ATRO	GTGA	ggt
	Consensus'	(040)	I CA.	L GC	G 'WI	CGCG	TA	GAG	CA	TTCC	GC AC	3 G
		10001									Sect	ion 17
		(689)	689		700	A SECTION CONTRACTOR	7	10		,720		73
	BCV N	(668)	CCC	TACCT	CTGG	TGTAA	CACC	TOAT	CATC	GCTG/	TCAA	TTC
0-4-m lmf47												
Avian infectious	DIONCHIAS N	(554)	CAGG	∔™ははかべ	CTAC	രമരനദ	CACC	STORY W.	78	Server services	-	25.0
	Consensus	(689)	C C	TGCCT	CTAG	TGTAA	ACC	TGA:	CATG	GCTG	TCAAZ	TTG
		(732)	732	7. TCTTGT TGTTGT	40		750		7	80		77
	BCV N	(711)	DAG	CRTCT	TOTAL	COASS	A Cilipar	HOAT	2 x 2 c c		a comparis	0.000
	MHV N	(714)	TGC	CPTOR	ካጥኮር	COTTO 2	COTO		2 2 2		ACT A	CARC
Avian infectious	bronchitis N	(594)	CCG	CCAGC	ΔΑΑΚ	ח השת מ	TONO	20 20 20		4100	20000	ال الواد
	Consensus	(732)	T G	CTTGT	ጥ ጥር	COMPA	On	SALES OF	-AGC	46	HOUR	rede.
						OULAN		GGC	MGG		Sect	
		(775)	775	.780		700						
	BCV N	(754)	COCC	780 AAGTA	F. 2482.44	,/90	D _ 9897	eren er	SOO		war needs	81
	MUVN	(757)	CALC	44.7	ACTA	AUCAG	ACUG	CCAZ	VAGA	AATC/	GACA-	- CA
Avian infectious	bronchitic M	(634)	AAA	AAGTA	ACGA	AGOAA	AGOG	CCA	MGA	AGTC/	GGCA-	-CA
Avian infectious	Concension	(034)	ACG	GCATT	AUTA	AGCAA	AAGG	CAG	VAGA	GATGO	CTCAT	'CGC
	Consensus	(775)	AAG	CAAGTA	ACTA	AGCAA	A TG	CCA	AGA	AATC	G CA	ĠA.
											Sect	ion 20
		(818)	818		,83	30		840		.85)	86
	BCV N	(795)	TAA	TTGAA	TAAD	ccccc	CCAG	AAGI	GGA	GCCCC	APAG	September 1970
Avian infectious												
	Consensus	(818)	PTAA	TTGAA	AAG	cccca	CCA	AAGE	ADD.	CC	AATTAA	DOD.
											- Secti	ion 21
		(861)	861		B70		880	•				
	BCV N	(838)	maria	CIGTI	200	Omoin	000	A A Lat	TELES !	890	2 49.745.10	90
	MHVN	(841)	mode	CAGTG	0000	4 6 7	4 4 T G	G G A	CAG	AUGCO	CCAAI	CAG
Avian infectious	hronchitie N	(720)	V CILL	Manage	CAGC.	wen Gil	TTTG	AAA	GAG	AGGCC	CCAAT	CAG
Avian Infectious	Consensus	(004)	MGTT	THIGH	CCCT	CGTAC	L'AAA	GGTA	AAG	AGGGI		3

		(00.4)		*						_ Section	22
	BCVA	(904)	904	910	10112 80	92	0	930			94
	MHIVA	(001)	ATT	TIGGT	GIGG	AGAA	TGTTA	AAACTT	GGÄACT	AGTGA	CC
Avian infactions	MITTY I	(004)	ATT	TTGGA	GCTC	TGAA	TGTTA	AAACTT	GGAACI	AGTGA	T.co
Avian Infectious											
	Consensus	(904)	ATT	TTGGT	GTG	GAA	TGTTA	AAACTT	GGAACT	AGTGA	FC
										Section	
		(947)	947			960		970			
	BCV N	(924)	ÁCA	GTTCCC	CATT	CTTG	17.0	- C2524	COCNOX	ocmos.	
A	MHA N	(927)	ACA	GTTCCC	CATT	CTTG	AGAGT	- TCGCA - TGGCT	OCA A CA	Compagn	1.0
Avian Infectious											
	Consensus	(947)	ACA	GTTCCC	CATT	CTTGC	AGAAC	T GCA		CCACAG	ŲĢ.
								1 GCA	CCARCA	GCTGGT	G
		(990)	990		1000		.1010			Section	
	BCV N	(966)	GTT	near recomm	diomin.	Tiela Sie	Section 15	AGPIEG	1020	219-1212	103
	MHV N	(969)	Critical	Chucana	TOMA.		ATH AG	AGUILGG AATI QQ	CAAAG	TGCAGA	A.
Avian infectious	bronchitis N										ιÀ.
	Consensus										
		(000)		111011	IGGA	TCTAG	ATTAG	ATTGG	CCAAAG	AGA	A
		(1033)	4000							Section:	25
	BCV N	(1000)	1033	104	40	.1	050	,1060)	1	07
	WHAV	(1009)	TiTG'	TCTEEG	AATC	TIGAC	GAGCC	1060 CAGAAC	GATGT	TEATOA	Ž.
Avian infectious	bronobitio At	(1009)	E	- Gregge	GGTG	CIGAT	GAACC	ACCAA	GATGT	GTATGA	GC
Avian infectious	Consensus			- 64 1	CAAC	CAGAT	GGGGT	PG 20	CTTAG	ATTTCA	ďΑ
	Consensus	(1033)	T	CTGGT	ATC	CTGAT	GAGCC	CC AA	GATGT	TATGA	Α'n
										- Section :	
		(1076)	1076			1090		.1100		- 1	118
	BCVN	(1052)	TEC	SCTATA	ATGG'	CGCAA	TTAGA	TTGACE	GTACA		
Arion infantia	MHVN	(1049)	TOC	ATATT	CAGG	IGCAG	TTAGA	TTGACE TTGAT	CTACT	CTACCT	7
Avian infectious	pronchitis N	(905)	TT-9	ACTACT	GT GG	recc-	-TAGAC	ATCACC	CGCAG	TOTOGAT	7 7
	Consensus	(1076)	TGC	ACTATT	TGG	FGCA	TTAGAT	TTGACA	GTAC	CAM CM	'can
										- Section 2	27
		(1119)	1119		.1130)	:1140	,			
	BCV N	(1095)	TTT	GAGAC	CAMA	-	telanda err	and divide the co	150 20 20 20 20 20 20 20 20 20 20 20 20 20		161
	MHV N	(1092)	BTTT	GAGAC	TATC.		TODAR	TGTTGA TGTTGA	ALGAG	AATTTG	ΑA
Avian Infectious	bronchitis N	(945)	TTA	GTANA	AARTT	rama [®]	TOTOM	TGIIGA	ATOAG	AATTTG	AΑ
	Consensus	(1119)	TTTT	GAGAC	Δm		TATE OF T	TGTTGA	TGGTG	ragg	Α̈́
						A	IGAA C	TGTTGA	ATGAG	AATTTG.	AA
	i	(1162)	1162	44	70		1100			Section 2	
	BCV N	(1134)	7007	ma mea		3.872	1180	,119	10	12	204
	MHVN	(1131)	100	LATCA	ACAAC	AAGA	IGGTAI	119 GATGA	ATATG	- GTCC	AΑ
Avian infectious	bronchitis N	(986)	2000	HACCA	AAGC	ATGG	regrec	-GATGA AGATGT	GG - TG7	- GCCC	ΑÀ
										CCCCC.	ÀΑ
	Consensus (102)	160	TACCA	A AAC	AAGG:	CGGTA	GATGA	A TGZ	GCCC	ת ת

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		(1205) 12	205		210			,122	20			1230)			124	7
		BCV N (1175) A.	ACC	AC	AG	CGT	CA-	G	dda	rgg	TCA	G	A	AGA	TOCA	545.7	÷
	Avian infectious	DIONCHIUS 14 (1020	JA.	TTO	ŊΑ	GA	CCT	GC-	-TA	CAZ	GA	ACA	AG-	TTC	TCC	GCGC	0000	î
		Consensus (1205) A	CC	AC	AA	CGT	A	G	CG	råg	ACA	GG	TA	GAP	GA	CAAC	
•													-				on 30	•
		(1248) 12	248				.126	30			1270			.1280			٠
		BCV N (1211) G	AGA	AΑ	AT	GAT	PAF	ATA	ÁGT	Circ	rar	AGC	ace	TAAA	Nacc	CTC	:
		BCV N (1211 MHV N (1214) A	TGA	AG	TA	GAT	AI	GUA	AGO	210	ron	222	ne e	2 2 2	7000	0.10.1	Ė
	Avian infectious	DIOLICITUS 14 (1009) A	CAA	CA	GC	CT C7	$\mathbf{A} \cdot \mathbf{A}$	ACD	10.726	ממני	מממ	or min	mry's	A MICH	- ninina		Ŀ
		Consensus (1248	Á	GA	AA		CAT	AT	ATA	AG.	CT.	rac	2 Z	CCC	A COM	AGGA	GGAN	
											٠		an	GCC	MAM		G1G1 on 31	
		(1291	۱ 12	91			.130	1		4	310			132	n	- Secu		
		BCV N (1254	í	TAC	CA	N'A	N 40 X 1	n la	rim S	7 20	1410	Arrive.	2. 5	102	100	.193.465	1333	
																	CATC	
	Avian infectious	bronchitis N (1111	, a	а те	22	CT.	CO A	4	CITA	GA		TTA.	400	CCAC	- 金	-GGA	TAGA	٠
		Consensus (1291	ía	736	ر با تقت	מ ת ת	ימידת	777	GW 2	j	Car	P. G.	A C C	TUE	AT G	AGGA	GWGC	ŕ
		(1201	,					·	GIM	GMC	MA.	ra G.	ACC	CAC	3A		AG	
		(1334	1 13	34		134	n		4	350			400			- Secti	on 32	
		BCV N (1294	1	- A-	100	Citi	- 4	12 5		200		5926 (42)	,136				1376	
		MHV N (1297	, A	200	-	Til	200	AA	CAA	GA	GGZ	111 (4)	Arana Starta			GĈCC	TATA	ļ
	Avian infectious	bronchitis N (1150	/ FPA		7 77	- G			34.1	CC	101	yn G	4.10	GCG:	CAGT	GCCA	GATC	
		Consensus (1334	1 24	a c.a.	wa.	GC.	ACAC	CIT	GUA	ATT	10.7	110	A.T.G	AAC	CCAA	GGTG		
			,	300		1		CA	GAA	1	T.G.	TG	ATG			GCC.	ATA	
		(1377	40	77												Section	on 33	
		DOM PL (1911	10	-	12.5	3797	20121	13	390	- 2.0	2277	,140	0			1414		
		BCV N (1325	01		- G	A A	ACI	CC	TCA	GAA	ATA	TA.	₩ - -					
	Avian infectious	MHV N (1340	G	T	AQ.	A A C	CATO	AC	TCT.	A A T	GTC	TA	Ŋ					
	Avian intectious	bronchitis N (1193	AC	MEG.	GG	GG	ATT	ÇA	GĊ₩	CTH	GGZ	GA	GAA	TGAC	TTG	TAA		
		Consensus (1377)	,	T	G.	AAC	JAT	CC	TCA	ΑT	GTF	TAL	A.					
		BCV N	SE	οт	n .	NTO .	: 98	83										
		MHV N					. 98											
,	Avian infectious b						99											

FIGURE 3B

Section '						
	30	20	,10	1	(1)	
ATGAGTAG	ATCCAAACATT	PTAGGTAAI	GGATGACGT	GATGT	(1)	HOBMPRO
ATGAGTAG					ı (1)	BCV M
ATGACTAG						MHV M Avian infectious brochitis virus M
						Avian infectious prochitis virus M Consensus
ATGAGTAG:					3 (1)	Consensus
Section 2			50	42	(42)	
8:		,60	DU	No William	(42)	HOBMPRO
PACTECTO:	TTTATATCTG	AGCACCAGI	TA OTICICA	CONTRACT	(10)	BCV M
JACTGGTG/	TTTACACCTG	AGCACCAGI	MACAUCA	80000	(10)	MHV M
TACGGCTG	TITATCAGTG	CAGCCTGT	LAGGERELA	Section Ca	(10)	Avian infectious brochitis virus M
SACTG-TG	TIGCACTOTT	CGGCAAAT	TEG CHACC	A AACT		Consensus
	STTTATCTCTG	AGCGCCAGT	TC CICCE	A AACI	(1-)	
Section 3		.100	90	83	(83)	
12:	,110 KGATGGAATT GAATGGAACT	100	CHAMPAN A THE	TOWN A KING	(80)	HOBMPRO
PTTCTTTG	GAATGGAATT	CCIAAAGG		TODAGO	(48)	BCV M
	MARIGGAACT	COTAAAGG	を	TICARO		
				ACAGTO	(36)	Avian infectious brochitis virus M
CTETCHCE	GARUGGAATI	TTOTAAGG	CAGTTGAGCT	ACAGTO	(30)	Avian infectious brochitis virus M Consensus
CTETCEC PATTTATA PTTCTTTG		TTOTAAGG	CAGTTGAGCT	ACAGTO	(30)	Avian infectious brochitis virus M Consensus
CTETCHOS CATTATAL PTTCTTTGG Section 4	GAATGGAATT GAATGGAATT	TTTTAAGG CCTTAAGG	CAGTTGAGCT	TGAGGC	(124)	Consensus
CTETCHOE CARTTATA PTTCTTTG Section 4	GAAUGGAATT GAATGAATT GAATGGAATT	TTTTAAGG CCTTAAGG	CAGTIGAGCU CTATTAAATT	TGAGGC	(124)	Consensus
CTETCHOE CARTTATA PTTCTTTG Section 4	GAAUGGAATT GAATGAATT GAATGGAATT	TTTTAAGG CCTTAAGG	CAGTIGAGCU CTATTAAATT	TGAGGC	(124)	Consensus
CTETCHOE CARTTATA PTTCTTTG Section 4	GAAUGGAATT GAATGAATT GAATGGAATT	TTTTAAGG CCTTAAGG	CAGTIGAGCU CTATTAAATT	TGAGGC	(124)	Consensus
CTETCECE FATTRATA FTTCTTTGC Section 4 16 EXATTTGC IAATTTGGT AGTTCGGT	TYANGGRATH THANGGRATH GALLANDER THANGGRA THANGGRANGE TANGGRANGE TANGGRANGE	140 TTACACACTACACTACACTACACTACACTACACTACAC	GAGT/GAGCT CTATTAAATT 130 ATACTACTT ATACTACTT ATACTACTT ATACTACTT TTCCTATTG	TGAGGC 124 GTATTA GTATTA GCATTA	(124) (121) (121) (89) (92) (77)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M
CTETCECE FATTRATA FTTCTTTGC Section 4 16 EXATTTGC IAATTTGGT AGTTCGGT	TYANGGRATH THANGGRATH GALLANDER THANGGRA THANGGRANGE TANGGRANGE TANGGRANGE	140 TTACACACTACACTACACTACACTACACTACACTACAC	GAGT/GAGCT CTATTAAATT 130 ATACTACTT ATACTACTT ATACTACTT ATACTACTT TTCCTATTG	TGAGGC 124 GTATTA GTATTA GCATTA	(124) (121) (121) (89) (92) (77)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M
CTETCHCE EATTTACA FTTCTTTACA Section 4 16 EATTTEC EATTTTEC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EATTTCC EATTTCC EATTTTCC EATTTTCC EATTTTCC EATTTTCC EAT	GAAUGGAATT GAATGAATT GAATGGAATT	140 TTACACACTACACTACACTACACTACACTACACTACAC	GAGT/GAGCT CTATTAAATT 130 ATACTACTT ATACTACTT ATACTACTT ATACTACTT TTCCTATTG	TGAGGC 124 GTATTA GTATTA GCATTA	(124) (121) (121) (89) (92) (77)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M
CTETCHCE FATTTATA PTTCTTTGC Section 4 16 ENATTTGC IAATTTGC ACTTCGCI CAGTTAGGC CAGTTTGGC Section 5	GARTATATT GARTGART 150 ANTGARTATT ANTGARTATT ANTGARTATT TATCATATT TATCATATT TATCATATT	TTTTTAAGG 140 TTATTACA TTATTACA TTATTACA TTATTACT	GAGTIGACTE CTATTAAATI 130 ATACJACTE ATACJACTE ATACJACTE ATACTACTE TTCCTATEG ATACTACTT	TGAGGC 124 GTATTA GEATTA CCATTA GTATTA	(124) (121) (121) (89) (92) (77) (124)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus
FCHETCHICE FATTTATA FTTCTTTGC Section 4 16- FATTTGC FATTTGC AATTTGC AATTTGC AGTATGC CAGTATGC Section 5 205	GARTATATT GARTATT GARTAGT 150 ANTGATATT ANTGATATT ANTGATATT TATCATATT TATCATATT TATCATATT TATCATATT TATCATATT TATCATATT TATCATATT TATCATATT TATCATATT TATCATATT	TTTTE AAG CCTTAAGG 140 TTATTAGA TTATTAGA TTCTTACT TTATTACT TTATTACT	GAGTIGACTO CTATTAAATI 130 ATACTACTO ETACTACTTI ATACTACTTI TTCCTATTO ATACTACTTI	TGAGGC 124 GTATTA GEATTA GCATTA GCATTA GTATTA 165 1 TATACA	(124) (121) (89) (92) (77) (124) (165) (162)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO
ECHERCHICE FATT TATAN Section 4 16 ENATT GET ASTREE ASTREE Section 5 20 LAGALTREE LOOK	GARTATER GARTATER GARTATER 150 ANTGATATE ANTGATATE ANTGATATER THE ACTA THE ACT THE ACTA THE A	TTTTEAGG 140 TTATTAGA TTATTAGA TTATTAGA TTATTAGA TTATTAGA TTATTAGT 180 ATGTTTGT	GAGTICAGO CTATTAAATI 130 ATACHACTEN ATACHACTEN ATTICTAGTIN TCCIATTOT ATACTACTTI	TGAGGC 124 GTATTA GCATTA GCATTA GCATTA GTATTA TATACA TATACA	(124) (121) (89) (92) (77) (124) (165) (162) (130)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M
FOREACHICA FATTOTTAGA Section 4 FRATTIGGO FRAT	GORATA GORATATATA GORATATATA 150 INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATATA	TTYERAGE 140 170 TATTAGE 170 TATTAGE 170 TATTAGE TTATTAGE TTATTAGE TTATTAGE TTATTAGE 180 ATGTTTGT	CAGTICAGCI CTATTAAATI 130 ATACTACTI ATACTACTI ATACTACTI ATACTACTI ATACTACTI ATACTACTI ATACTACTI ATACTACTI ATACTACTI ATACTACT ATACTACT ATACTACT AGTICACAGI AGTICACAGI AGTICACAGI	TGAGGC 124 GTATTA GEATTA GCATTA GTATTA 165 TATACA	(124) (121) (89) (92) (77) (124) (165) (162) (133)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M
ECHETCHICE EATTTA TATA FITTCTTTGC Section 4 16 ENATTTGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTIGC	GENERAL STANDARD STAN	140 140 174716A 174716A 174716A 174716A 174717A 180 174717A 174717A 174717A	CACTLEGGO CTATTAATT 130 ATALTACTH THICKETACTH THICKETACTH THICKETACTH 170 ATACTACTH 170 ACTACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH	TGAGGC 124 GTATTA GCATTA GCATTA 165 TATACA TATA	(124) (121) (89) (92) (77) (124) (165) (162) (130) (133) (118)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M
ECHETCHICE EATTTA TATA FITTCTTTGC Section 4 16 ENATTTGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTIGC	GENERAL STANDARD STAN	140 140 174716A 174716A 174716A 174716A 174717A 180 174717A 174717A 174717A	CACTLEGGO CTATTAATT 130 ATALTACTH THICKETACTH THICKETACTH THICKETACTH 170 ATACTACTH 170 ACTACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH	TGAGGC 124 GTATTA GCATTA GCATTA 165 TATACA TATA	(124) (121) (89) (92) (77) (124) (165) (162) (130) (133) (118)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M
ECHETCHICE EATITA MAA FITTCTTTGC Section 4 16 ENATITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTITGC ANTIGC A	GORATA GORATATATA GORATATATA 150 INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATA INTEGRIATATATA	140 140 174716A 174716A 174716A 174716A 174717A 180 174717A 174717A 174717A	CACTLEGGO CTATTAATT 130 ATALTACTH THICKETACTH THICKETACTH THICKETACTH 170 ATACTACTH 170 ACTACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH TACTACTH	TGAGGC 124 GTATTA GTATTA CCGCTAT GTATTA GTATTA TATACA TATACA TATACA TATACA TATACA TATACA TATACA	(124) (121) (89) (92) (77) (124) (165) (162) (130) (133) (118) (165)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus
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ECHETCHICE ATTTA TATA EATTTA TATA 16 ENATTTECH A TTTCGET ACTTA TOGA Section 5 200 AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT AAGATCAT Section 6	GARTAGERATE GARTGARTAGER ANT CATACHT ANT CATACHT ANT CATACHT THE CATACHT TATCATATTG THE TATCATATTG THE TATCATATTG THE TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATTT TATCATATAT TATCATATAT TATC	140 140 140 140 140 140 140 140 140 140	CACTTAGGO CTATTAGATT 130 ATACTACTTAG ATACTACTTAG ATACTACTTA ATACTACTTA 170 AGT CGGGT AGACAGGAGT AACAAGGAGT AACAAGGAGT	TGAGGC 124 124 124 124 124 124 124 124 124 12	(124) (121) (89) (92) (77) (124) (165) (162) (133) (118) (165) (206) (203)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO
TELETONICE ATTITATA TETTOTTTG Section 4 16 ENATTITGG ASTITGG Section 5 20 ANATTIGG ANATOM ANATOM ANATOM ANATOM ANATOM ANATOM ANATOM ANATOM ANATOM Section 6 26 ENATOM ANATOM ANATOM ANATOM ANATOM ANATOM Section 6 26 ENAME ENTRACE	GENERAL GENERA	TTY MAAG CCTTAAGG 140 1TTATTACA TTATTACA TTATTACA TTATTACT 180 ATGTTTGT ATGTTTGT ATGTTTGT ATGTTTGT ATGTTTGT ATGTTTGT ATGTTTGT	CACTTO GOOD 130 130 ATACTACTA ATACTACTA ATACTACTA ATACTACTA ATACTACTA ATACTACTA ATACTACTA ATACTACTA ATACTACTA ATACTACTA AAGTCGGAST AAGTCGGAST AAGTCGGAST AGTCGGA	TGAGGC 124 GTATTA GCATTA GCATTA 165 TATACA TATACA TATACA TATACA TATACA TATACA TATACA TATACA TATACA TATACA CO TATACA	(124) (121) (89) (92) (77) (124) (165) (162) (130) (133) (118) (165) (206) (203) (203) (171)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M HOBMPRO BCV M
ECHETCHIC EATT TA MAZE TO THE COLOR OF THE C	GARTAGE AND THE CONTROL OF THE CATACTA THE	140 140 140 140 140 140 140 140 140 140	CACTAGACT 130 ATACHACTE ATACHACTE ATACHACTE ATACHACTE ATACHACTE ATACHACTE 170 AAGT CGCACT AACAACGAG AACAACGAG AACAACGAG TEGGAGT TE	TGAGGC TGAGGC TGAGGCATTA TATACC TTATAC	(124) (121) (89) (92) (77) (124) (165) (162) (130) (133) (118) (165) (206) (203) (171)	HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Consensus HOBMPRO BCV M CONSENSUS HOBMPRO BCV M MHV M CONSENSUS
ECHETCHICE ATTTATATA FITCTTTG Section 4 AATTTGGA AATTTGGA AATTTGGA AGATTGGA AGATTGGA AGATTGGA AGATTGGA AGATTGGA AGATTGGA AGATTGAA	GARTAGE AND THE CONTROL OF THE CATACTA THE	TTYMEAG CCTTAAGG 140 TTATTAGA TTATTAGA TTATTAGT TTATTAGT 180 ATGTTTGT ATGTTTGT ATGTTTGT ATGTTTGT ATGTTTGT CCGTTMGA ATGTTTGT CGGTTGT CGGTGGCCG GTGGCCG GTGGCCG GTGGCCG	CACTTAGACT 130 ATACTACTACT ATACTACTACTACTACTACTACTACTACTACTACTACTAC	124 GTATTA GCATTA GCATTA GCATTA GCATTA GTATTA 165 1. TATACA TATACA TATACA 206 CATTATTA CATTTATACA CATTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTATACA CATTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTATACA CATTTACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTATACA CATTTACA CATTTATACA CATTTACA CATTTACA CATTTACA CATTTATACA CATTTACA CAT	(124) (121) (89) (92) (77) (124) (165) (162) (130) (133) (118) (165) (206) (203) (171) (174) (159)	HOBMPRO BCV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M Consensus HOBMPRO BCV M MHV M Avian infectious brochitis virus M

Section 7			
287	270	260	(247) 247
ATGTGTATCTTGGC	CGCATTGAATA	TCAATTGCGTATAC	HOBMPRO (244) TT
A TICTO TO A TO COMPOSE	TOTALIZATION	TCAATEGCCTCTAT	BCV M (212) TT
A TOTAL TO A MANUAL COMPA	アはつだってカカカカヤカ	TTAACTCCCTCTAT	MHV M (215) TT
CARGAGGTCTTTTT	TCGACCAAATA	TTTCATGTATATAT	Avian intectious procnitis virus M (200) TA
ATGTGTATCTTGGC	TGCGTTGAATA	TTAATTGCGTATAT	Consensus (247) TT
Section 8			
328	310	300	(288) 288
a hom almon a michigan	CCATAGTGGC	TCTATAGTTTTTAC	HOBMPRO (285) C
TTTAMOARRICANT	CHATACHECCE	TCTATAGTTTTTCAC	BCV M (253) 宝宝
A THE A PLANT OF THE PARTY OF THE A	CTATACTOTO	DCTABAGTGTTTAC	MHV M (256) ##
TATOMATATIONAL	CAGMCCATTCAC	GUGAAAATACETAC	Avian intectious brochius virus M (241) GC
ATTAT ATCTCCAT	CTATAGTGGCC	TCTATAGTTTTAC	Consensus (288) TT
Section 9			
369	350	340	(329) 329
demmin a marida a dime	NORA TO A COM	CTATTOTOTON	HOBMPRO (326) 75
Commanus of a day	ACTATORISCTT	etatttlöignata	BCV M (294) TG
Craminacon	ACCAMC ROOM	GTATTTTGTTAATA	MHV M (297) TA
		2000 1000 1000 1000 1000 1000 1000 1000	Auton Infantious buschille ulura \$4 (000) = 2
CHETTALCOCOCO	7 6 T 7 T C A C 7 C A	ATATTCCATTCACA	Avian intectious prochitis virus ivi (282) AG
CTTTTAAGCGGGGGG	AGTATOAGACII	ATATEGGATECAGA GTATTTTCTTAATA	Avian Infectious brochitis virus M (282) Ac Consensus (329) TG
CTTTAAGCGGGGC GTTTATTAGGACTG	AGTATCAGACI AGTATCAGGT1	ATATEGGATMCAGA GTATTTTGTTAATA	Consensus (329) TG
CTTTAAGCGGGTC GTTTATTAGGACTC Section 10	AGTATCAGGT1	GTATTTTGTTAATA 380	(370) 370
CTTTAAGCGGGTC GTTTATTAGGACTC ———————————————————————————————————	AGTATCAGGT1	GTATTTTGTTAATA 380	(370) 370
CTTTAAGCGGGGTC GTTTATTAGGACTC ———————————————————————————————————	AGTATCAGGT1 390 CAACGGAGAAA	GTATTTTGTTAATA 380 GTTTTTGGAGTTBG	(370) 370 HOBMPRO (367) GA BCV M (335) GA
CTTTTAGCGGGGTC GTTTATTAGGACTC —— Section 10 400 411 CAAACAACTTGATC CAAAGAACTTGATC	AGTATCAGGT1 390 CANCOCAGAA	380 GTTTTTGGAGTING GTTTTTGGAGTING	(370) 370 HOBMPRO (367) GN BCV M (336) GC
CTTTTAGCGGGGTC GTTTATTAGGACTC —— Section 10 400 411 CAAACAACTTGATC CAAAGAACTTGATC	AGTATCAGGT1 390 CANCOCAGAA	380 GTTTTTGGAGTING GTTTTTGGAGTING	(370) 370 HOBMPRO (367) GN BCV M (336) GC
CTTTAAGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	390 CANCONGANA CANCOCAGNAN CANCOCAGNAN CANCOCCIGAN TANCOCTIGAGT	380 GTTTTTGGAGTTTGGAGTTTGGAGTTTGGAGTTTGGAGTTTGGAGTTTGGAGTTTGGAGTTTGGAGGTTTGGAGGA	Consensus (329) TG (370) 37(HOBMPRO (367) GA BCV M (335) GA MHV M (338) GC Avian Infectious brochility virus M (323) GG
CTTWAAGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	390 CANCONGANA CANCOCAGNAN CANCOCAGNAN CANCOCCIGAN TANCOCTIGAGT	380 GTTTTTGGAGTING GTTTTTGGAGTING	Consensus (329) TG (370) 37(HOBMPRO (367) GA BCV M (335) GA MHV M (338) GC Avian Infectious brochility virus M (323) GG
CTTTAT GGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AGTATCAGGTT 390 CANCOCONA CANGOCOSA TANCOCOSA TANCOCOSA CANCOCOSA CANCOCOSA TANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOS CA	380 GTATTTTGATAATA GTATTTGAGTATC GTTGGTGGAGTATC GCTGGTGGAGTATC GCTGGTGGAGTATC GTTGGTGGAGTTTC	(370) 370 HOBMPRO (367) GA BCV M (335) GA MHV M (338) GC Avian infectious brochilits virus M (323) GG Consensus (370) GA
CTTTATTAGGAGACTC GTTTATTAGGACTC A00 41 CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGAT CAAACAACTTGATC CAAACAACTTGAT Section 11	AGTATCAGGTT 390 CANCOCONA CANGOCOSA TANCOCOSA TANCOCOSA CANCOCOSA CANCOCOSA TANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOSA CANCOCOS CA	380 GTATTTTGATAATA GTATTTGAGTATC GTTGGTGGAGTATC GCTGGTGGAGTATC GCTGGTGGAGTATC GTTGGTGGAGTTTC	(370) 370 HOBMPRO (367) GA BCV M (335) GA MHV M (338) GC Avian infectious brochilits virus M (323) GG Consensus (370) GA
CTMEAR COGGORE GTTTATTAGACTC Section 10 A00 41 CAAACAACTTGAT CHAAACAACTTAT CTAAACAACTTGAT CTAAACAACTTGAT CTAAACAACTTGAT 40 640 640	AGTATCAGGTT 390 CANCODAGAA CANGOCAGAA TAACOCAGAA CAACCCAGAAA 430 GAACCAGAA	GTATTTTGTTAATA 380 GTTTTTGGAGTTTC GTTGGTGGAGTTTC BCTGGTGGAGTTTC A20 AATGGTGGAGTTTC A20 ATGGTAGAGGTTC	(370) 370 HOBMPRO (367) GA BOV M (335) GA MHV M (338) GC Avian infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (408) TG
CTTTATTAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AGTATCAGGTT SOC STATE OF STAT	GTATTTTGTTAATA 380 GTTTTTGGAGTTTC GTTGGTGGAGTTTC GCTGGTGGAGTTTC GTTGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG	(411) 411 HOBMPRO (406) TG AVIan Infectious brochilits virus M (323) GC Consensus (370) GA (411) 411 HOBMPRO (406) TG BCV M (376) TG MHV M (376) TG
CTTTATTAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AGTATCAGGTT SOC STATE OF STAT	GTATTTTGTTAATA 380 GTTTTTGGAGTTTC GTTGGTGGAGTTTC GCTGGTGGAGTTTC GTTGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG	(411) 411 HOBMPRO (406) TG AVIan Infectious brochilits virus M (323) GC Consensus (370) GA (411) 411 HOBMPRO (406) TG BCV M (376) TG MHV M (376) TG
CTEMAR CONGROUNT CTTATTAGACT A00 411 CAAACAACTTGAT CHAA CAACTTGAT CTAA CAACTTGAT CTAA CAACTTGAT 40 45 GTINGGCCGATAAI GETAGACCGATAAI GETAGACCGATAAI GETAGACCGATAAI GETAGACCGATAAI	AGTATCAGGTT S90 CANCECAGAA GARCECGAAA TAAGGTGAAA A30 BAACAATGTAT SAAGGATGTAT GRACTGTGTTAT	GTATTTTGTTAATA GTTTTTGGAGTATC GTTGGTGGAGTATC GCTGGTGGAGTATC ATGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG	Consensus (329) TG (370) 37C HOBMPRO (367) GA BCV M (336) GA MHY M (338) GC Avian infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (406) TG BCV M (376) TG MHY M (379) TG Avian infectious brochitis virus M (359) T-
CTTTATTAGGACTC A00 411 CAACAACTTGATC CTAACAACTTGATC CTAACACTTGATC CAAACACTTGATC CAAACACTTGATC CAAACACTTGATC CAAACACTTGATC A40 Section 11 A40 CTTAGGCCATTAAACTTGATC CTTAGGCCATTAAACTTGATC CTTAGGCCATTAAACTTGATCATCATCATCATCATCATCATCATCATCATCATCATCA	AGTATCAGGTT S90 CANCECAGAA GARCECGAAA TAAGGTGAAA A30 BAACAATGTAT SAAGGATGTAT GRACTGTGTTAT	GTATTTTGTTAATA 380 GTTTTTGGAGTTTC GTTGGTGGAGTTTC GCTGGTGGAGTTTC GTTGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG	Consensus (329) TG (370) 37C HOBMPRO (367) GA BCV M (336) GA MHY M (338) GC Avian infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (406) TG BCV M (376) TG MHY M (379) TG Avian infectious brochitis virus M (359) T-
CTTTATTAGGACTC A00 A00 CTTTATTAGGACTC A00 CTAACAACTTGATT CTAACAACTTGATT CAAACAACTTGATT CAAACAACTTGATT A40 Section 11 A40 GTTTAGGCCGATTAAACTTGATC GTTTAGGCCGATTAAACTTGATC GTTTAGGCCGATTAAACTTGATC GTTTAGGCCGATTAAACTTGATCAACAACTTGATCAACAACAACACAACAACAACAACAACAACAACAACAA	AGTATCAGGTT 390 CANCESTAGA CARGERAGE GARCECGAAA TAACOCTAGAA 430 GARCANTGTAT TAACOCTAGAA CARCANTGTAT TAACOCTAGAAA GARCANTGTAT TAACGATTAT TAACGATTAT TAACGATTATAT	GTATTTTGTTAATA 380 GTTTTTGGAGTATC GTTGGTGAGTATC GCTGGTGAGCTTC AATGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATAGATATGAAAGG	(452) TG (370) 37(HOBMPRO (367) GA BCV M (335) GA MHV M (338) GC Avian Infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (408) TG BCV M (376) TG MHV M (379) TG Avian infectious brochitis virus M (359) TG Consensus (411) TG
CTTTATAGGACTI A00 A00 CTAACAACTAGTAT CTAACAACTAGTAT CTAACAACTAGTAT CTAACAACTAGTAT CTAACACTAGTAT CAAACTAGTAT CAAACTAGTAT A40 Section 11 A40 GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA GTTTAGGCCGATTAA Section 12	AGTATCAGGTT 390 CANCESTAGA CARGERAGE GARCECGAAA TAACOCTAGAA 430 GARCANTGTAT TAACOCTAGAA CARCANTGTAT TAACOCTAGAAA GARCANTGTAT TAACGATTAT TAACGATTAT TAACGATTATAT	GTATTTTGTTAATA 380 GTTTTTGGAGTATC GTTGGTGAGTATC GCTGGTGAGCTTC AATGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATAGATATGAAAGG	(452) TG (370) 37(HOBMPRO (367) GA BCV M (335) GA MHV M (338) GC Avian Infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (408) TG BCV M (376) TG MHV M (379) TG Avian infectious brochitis virus M (359) TG Consensus (411) TG
CTTTATAGGCGGCGTGGCTTTATTAGGCGGTAATCCAAACACTTGATCAAACACTTGATCAAACACTTGATCAAACACACTTGATCAAACACACTTGATCAAACACACTTGATCAAACACACAC	AGTATCAGGTT 390 CAAGUCAGAAA CAAGUCAGAAA CAAGUCAGAAA A30 BAACAATGTAI GAAGGATGTAI GAAGGATGTAI GAAGGATGTAI GAAGGATGTAI GTACTATGTAI A70 CTGAGGGTCAC	GTATTTGTTAATA 380 GTTTTGGGGTTEC GTTGGTGGGGTTTC AATGGTGGGGTTTC A20 ATACATATGAAGG ATACATATGAAGG -AGGT-TCAATAC ATAGATATGAAGG -AGGT-TCAATAC ATAGATATGAAGG ATAGATATGAAGG AGGT-TCAATAC AGGT-TCAA	(370) 370 HOBMPRO (367) 370 BCV M (335) 62 MHV M (338) 62 Avian Infectious brochitis virus M (323) 66 Consensus (370) GA (411) 411 HOBMPRO (408) 36 BCV M (376) 76 MHV M (376) 76 Avian infectious brochitis virus M (359) T- Consensus (411) T6 (452) 452 HOBMPRO (449) 76
CTILLAG CGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AGTATCAGGTT 390 CANCERAGA CANCECTAGAA GARCECGAAA 430 GARCANTGTAT FRAGGATCTAT GTRACTATTATATATATATATATATATATATATATATATAT	GTATTTTGTTAATA 380 GTTTTTGGAGTATC GTTGGTGGAGTATC GCTGGTGGAGTATC GTTGGTGGAGTTTC A20 ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATACATATGAAAGG ATAGATATGAAAGG AGGATATCATACTC	Consensus (329) TG (370) 37C HOBMPRO (367) GA BCV M (335) GA MHY M (338) GC Avian infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (408) TG BCV M (376) TG MHY M (379) TG Consensus (411) TG Avian infectious brochitis virus M (359) TG Consensus (411) TG HOBMPRO (449) TG BCV M (417) TG
CTTTATTAGGACTO A00 410 CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC CAAACAACTTGATC GAAACAACTTGATC GTTAGGCCGATAAT GTTAGGCCGATAAT GTTAGGCCGATAAT GTTAGGCCGATAAT GTTAGGCCGATAAT GTTAGGCGGTAAT A80 ACAACAACAACAACAACAACAACAACAACAACAACAACA	AGTATCAGGTT 390 CANCESTANA CANCECTORA MACCCCORA ASO CANCECTORA ASO CANCERTA TORACOLTORA ASO CANCESTATORA GTACTAT GTACTAT TACGGTCAC CTTACGGTCAC CTTACGGTCAC	GTATTTTGTTAATA 380 GTTTTTGGGGTTTGGGGTTTGGGGGGGGGGGGGGGGG	Consensus (329) TG (370) 37C HOBMPRO (367) GA BCV M (335) GA MHY M (338) GC Avian infectious brochitis virus M (323) GG Consensus (370) GA (411) 411 HOBMPRO (408) TG BCV M (376) TG MHY M (379) TG Consensus (411) TG Avian infectious brochitis virus M (359) TG Consensus (411) TG HOBMPRO (449) TG BCV M (417) TG

						Section 1
(493) HORMPRO (400)	493	500		510	,520	53
HOBMPRO (490) BCV M (458)	ATOT	TTACAT	TCAAGGT7	TAAAAC	TAGGTACT	GGCTATTC
Auton Infantiary based in	ACCT	CTATAT	GEAAGGTO	TTAAGC	TAGGCACT	GGCTATTC GGCTTCTC
Consensus (493)	ATCT	TTACAT	CAAGGT	TAAAGO	TACCUACU	CITIATIG
		-			INGGINCI	
(534)	534	.540		^		Section 1
HORMORO (534)	Tricicioni	- WWW.	55	0	560	57
HOBMPRO (531) BCV M (499)	1.666	GAGATT	receage:	TATATG	ACTGTT GC	TAAGGTTA
MHV M (502) Avian infectious brochitis virus M (460)	TTGT	CTGATT	CCCTGOT	TATGET	CACTEGE	Thheemem
Consensus (534)	TTGT	CTGATT	CCCTCCT	TATCTC	CTCTTCAA	CCAGACGA
					10101160	IMAGGTUT
(575)	575	580	590			Section 1
HORMPRO (572)	3 (400) ct /	LANCE TO SERVICE	7200	0.0 (4) (2)	,600	61
HOBMPRO (572) BCV M (540)	ACACI	- Terreca	AATATAA	GEGTGO	TTTCTTG.	ACAGGATA
MHV M (543) Avian infectious brochitis virus M (496)	TCACC	TTTGC	CTTATAA	GGGGGC	TTCTTAN	CANCOTA
Consensus (575)	TCACC	CTGTGC	CATATAA	CCGTCGT	יסטטטפייייייייייייייייייייייייייייייייי	
(616)	616		630	-		- Section 16
HOBMPRO (613) BCV M (581)	GCCNA	A Amazon	530	Carlot Acceptance	40	65
BCV M (581)	COCONI	ALTAG	COLLTING	CTGTTTT	CTGTTAAG	CCAVAGT
MHV M (584) Avian infectious brochitis virus M (531)	ACGGT	GTTAGO	GGTTTTG	CTGTTT	TGTGAAG	CCAACGT
Consensus (616)	GCGAI	ACTAGI	GGTTTTG	CTGTTTA	TGTTAAGT	ירים אמרים
						- Section 17
(657)	657		.670	,680	`	•
HOBMPRO (654)	CCTAA	TTACCC	A COMPANIE	The state of the s	THE PERSON NAMED IN COLUMN	69
BCV M (622) MHV M (625)	GGMAN	Tracco	ACTOUR	TCAACGC	AAAAGGGT	TCTGGCAT
MHV M (625)	000		ACTRECT	TCAACCC	AAAAGGG1	TCTGGCAT
MHV M (625) Avian infectious brochitis virus M (562)	COMMA	LIACUG	ACTGOCC	TCAAATA	AACCGAGT	GGCAG
Consensus (657)	GGTAA	TTACCG	ACTGCCA:	TCAACCC	AAAAGGGT	TCTGGCAT
The state of the s						- Section 18
(698)	698		,710	720		
HOBMPRO (695)	GGACA	CCCCAT	TOWNER	2 x m - 17 2	D 125	738
						TTTAAGGA
MHV M (663)	CCACA	COGCAT	TOTICAC	AATAAT	ATCTAA	
wlan infectious brochitis virus M (601)	GGAGA	CUGUAL	TGTTGAG	9;	ATCTAA'	
Consensus (698)	GGACA	CCGCAT	TGTTGAG	TAATAAF	ATCTAA	
/700					Se	ction 19
(739) HOBMPRO (736)						
FOURTON (736)		Q ID NO:				•
BCV M (694)		Q ID NO:				
MHV M (688) Avian infectious brochitis virus M (601)		Q ID NO:				
(601)	SE	ON DI O	9904			
Consensus (739)						

FIGURE 3C

											Section 1
	(1)	1	.10			20		,30	40		50
HOBHEGA	(1)	CTAA	ACTCAG	TGAA	ATGT	rtttg:	TTC	TAGAT	CTATT	TACTTA	GCTGCAT
BCV HE	. (1)	CTAAT	ACTOAG	TGAA	ATGT	TTTTG	mr ca	TAGAT	PTGTTO	TACTTA	SCIBCAT
MHV HE	(1)	130 to 1377.	SA WARRA	**************************************	1939	45.5.5.5.	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	CO. Mara S. C.	SICHER PR	4377777	62. 6.34
	- 22	00333	00000	mas s :		mmmma	amma	magame		ma coma	GCTGCAT
Consensus	(1)	CTAA	AUTCAG	TGAA	AATGT	TTTTG	JTTC	TAGAT	rr Tre	TAGTTA	
											Section 2
	(54)	54	60		,70		,80		90		100
HOBHEGA	(54)	AATTO	GTAGC	TTAG	STTTT	TACAA	COCT	CTACC	AATGIT	GTTTCG	CATGITAL
BCV HE	(54)	AATT	GTAGC	CTAG	STTTT	GACAA	TOCT.	ETAGC.	AATGTT	GTTTCC	CATTTAI
MHV HE	(1)										
Consensus	(54)	DATTO	CCTACC	TAGG	մահահան 1	ACAA	CCTC	CTACC	TAPGTT	GTTTCG	CAT TA
00110011000	(0.,										- Section 3
	(407)	407		.12	n .	,13		.14	0		15
	(107)	TO/	DOMESTIC STREET							and the same	
HOBHEGA	(107)	ATGG	ACATTG	GTTT	TTATT	TGUTE	ACAG	CGFEC	AGATTG	TAATOR	TATTCT
BCV HE		ATGG	ACATTC	GTTT	TTATT	TGGTG	ACAG'	regrie	AGATTG	TAATC	TGTTGT
MHV HE	(1)										
Consensus	(107)	ATGG	AGATTG	GTTT'	TTATT	TGGTG.	ACAG:	rcgttc.	AGATTG	TAATCA	TTGT
											— Section 4
	(160)	160		,170		.180		.190		200	21
HOBHEGA					TAA TOT	ATTICT	TATA	recare	TTAATC	CTETTO	TGTGTG
BCV HE											TGTGTG
MHV HE	(1)			29,90	tipe to	Co. Fara	A STANTA	e de la companya della companya de la companya dell	t anna anna	ng a ga	HOW AND THE
MU Á UE	(400)						m > m > .	TGGACC			TGTGTG
Consensus	(100)	A IA	CAACC	CCC	IAMLI	MITCI	TWIW	IGGMCC	TIMMIC	.CIG .	— Section f
								_	250		
	(213)	213	220	100000	230		24		250	100 20 111 FO 402 FO	26
HOBHEGA	(213)	PTCT	ggtalaa	ATAT	CATCT	AAAGC	TOGC	AACTCO	ATTTTT	AGGAG	DADTTU
BCV HE	(213)	THET	GGTAAA	ATAT	CATCT	AAAGO	TGGC.	AACTCC	APTTTT	AGGAG	TTTTCAC
MHV HE	(1)										
Consensus	(213)	TTCT	GGTAAA	ATAT	CATCT	AAAGC	TGGC	AACTCC	ATTTTT	AGGAG	DADTTT
											Section 6
	(266)	266			280		290		300		31
HOBHEGA	(200)	Bass	diria mun	mer o'm	2 2 mm 2	02.02.5	And A			2012 20 00 00 5	PTTATGA
	(200)	200	TOTAL LA	ALA		CACAC		AGG 1.CA		100	
BCV HE			WGA LUI	TAT WIT	MATTA	CACAG	A POPA	HOUT CA	WARNEY!	正。据证证 价	CTATGA
MHV HE	(1)										
Consensus	(266)	TTAC	CGATTI	TTAT	AATTA	CACAG	GCGA	AGGTCA	ACAAAT	TATTT:	r TATGA
											— Section `
	1010	319		,330		340		350		360	37
	(319)										
HOBHEGA	(319)	GGTC	TTAATT	TTAC	GCCTT	ATCAT	GCCT	TTAAAT	GCAAC	CGTTCT	GTAGTA
HOBHEGA BCV HE	(319)	GGTG	TTAAT	TTAC	GCCTI	ATCAT	GCCT	TTAAAT TTAAAT	GCACC	CGTTCT	GTAGTA
BCV HE	(319)	GGTG GGTG	TTAATI PTAATI	TTAC TTAC	GCCT1 GCCT1	ATCAT ATCAT	GCCT	TAAAT TAAAT	GCACCI GCACCI	CTTCT CTTCT	GTAGTA GTAGTA
	(319 (319 (1	GGTG GGTG	PTAATI)A‡E	GCCTT	A#CA1	GCCT	TTAAAT TTAAAT TTAAAT	gcacci	Chich	GTAGTA GTAGTA

	(372)	372		380			390		.400	<u> </u>					ion 8
HOBHEGA	(372)	TG	TATT	ידיכים	mara	CAAR	D 2 2	e Sale		7.77	20. 10.	410	-	TAAGA	424
BCV HE	(372)	TGA	TATT	TEGA	TOCA	CARA	0.7. 5. 2	000	TIGIN	11.0	ACTO	AGG	TTA	TAAGA TAAGA	ATA
MHV HE			an said d												
Consensus		TCD	ייים מיים	יייים מיייי	1,000 1,000	CAA	AAA	GCT	CGATT	TTA	GCCC	GAG	rgta	TGAGA	AGA
	(0.2)	102	TEST	LGGA	TGCA	GAAT	LAAA	GGC	PTGTT	TTAT	ACTO	'AGG'	LLL	TGAGA TAAGA	ATA
	(425)													Sect	ion 9
HOBHEGA	(420)	420	430	J 	EAST TO A	440			150		460				477
BCV HE	(420)	166	F 1.6 T	GTAT	CGICA	GCCI	TAC	TTT	rgtra	ATG	ACCA	TAT	TTT	ATAAT	CCC
MHVHE															
Consensus															
00113611303	(423)	166	CTGT	GTAT	CGCA	GCCI	TAC	TTTT	CTTA	ATG	ACCA	TAT	TTT	ATGGA ATAAT	GGC
														- Sectio	n 10
	(478)	478			490			500		510)		520		
HOBHEGA	(478)	TCT	GCAC	AATC.	PACA	COTO	TTT	GTAZ	ATCT	GGT	Chien	DO P	1000	AATAA	
BCV HE															
MHVHE	(97)	AAT	GCAA	AGCC	CGCC	TCCA	TIT	GCA	AGAC	AATT	Crimin	200	na.c	aataa Aataa	
Consensus	(478)	TCT	GCAC	AATC:	FACA	GCCC	TTT	GTAZ	ATCT	GGTA	CTTT	MGT.	CIPIT	AATAA AATAA	and.
											.0111	MGI	C11.	– Sectio	CCC
	(531)	531		.540			550		.56	30		570			
HOBHEGA	(531)	TGC	ATAT	ATAG	Tec	BC A.S	Frein	מה כח	A 7145	4.000	en witer en	PATER A	ALE CANCE	gerrd	583
BCV HE															
MHV HE															
Consensus	(531)	TGC	ATAT	ATAG	י יישי	TCDD	CCT	2 2 1111	widt	THAT	研究技術	ACT.	TGA	AGTG GTTG	AGG
									1.66	GGAT	TATT	ATTA	TAA	GTTG.	AAG
	(584)	584	.59	90		600			610					_ Section	
HOBHEGA	(584)	CTG	ATTE	PPARE	er in	CACC	mma	north c	-2 - CELL	E-LFE-SE	620	CONTRACTOR OF	,	itatt	636
BCV HE	(584)	CTC	E dimini	mm a m	200	0200	TILG	LUAC	GAGT	ATAT	CGTA	COAC	TAT:	TATT.	TTT
MHV HE	(203)	CTA	A TT T	CACAC	777	7700	116	Leve	GAGI	ATAT	CGTA	CCA'C	TTT T	TATT TATT TGGT	TTC
Consensus	(584)	CTG	ATTT	TT A TT	TO COL	77.77	TIG	LOAT	GAAT	TTAT	AGTA	CCGC	TCT	TGGT	TTT
	• •				. 1 61 (CMGG	TIG	FGAC	GAGT	ATAT	CGTA	CCAC	TTTC	TATT	TTT"
	(637)				650									 Section 	า 13
HOBHEGA	(637)	70 72 T	and N	a Calabia	<u>000</u>			660		6	70				689
BCV HE	(637)	4	360.	TTTOA	T.KG.			TC	GAAT	ACA-		-AAG	TATI	ATGA	rga.
Consensus	(637)	WW T		AGTTI	AAG	JGCA	GCT	CTTC	GGAT	CTG	CCAA	FAAA	TATI	ATGA'	PGA
00110011003	(031)	MAC	GGCAI	AGTTT	TTG			TC	GAAT	ACA		AAG	TATT	ATGA	rga.
	(000)													Section	
HORUGOA	(690)	690		,700)		,710		7	720		.730			
HOBHEGA	(6/5)	TAG:	rca a i	PATTA	TTTI	CAAT	AAAC	JAÇA	CTGG	CTT.	ATTE	F-12-74-7		AATT	1/1° %
Consensus	(690)	TAG:	rcaa'ı	CATTA	TTTI	TAAT	AAAC	ACA	CTGGT	CTT	4.44	TCC	O T T	AATT	υA

													Section	15
	(743)	743	,750			60		770		,780				795
HOBHEGA	(728)	e			AACCA									
BCV HE	(728)	C		-TGA	AAOCA	TTAC	ZA						TTGT	
MHV HE	(362)	CCTT	GATG	TTGG	CAACA	CTGC'	FAAGG	ATCC	GGĞŢ	CTIG	ATCT	CAC,	TŢGC	\GG
Consensus	(743)				AACCA			Č	TGGT	TTTG	ATCT	TĂA'	TTGT	ΑT
						·							Section	16
	(796)	796			810		820			30				848
HOBHEGA					CCTCI									
BCV HE					CCTCT									
MHV HE	(415)	TATC	TTGCA	TTGA	CTCCT	GGTA	ATTAT	AAGC	OTG	GTCC	TTAG	AAT	ATTT	3T I
Consensus	(796)	TATT	TAGTI	TTAC	CCTCT	GGTA.	PATTA	TTAC	CCAT	TTCA	AATG			
													Section	
	(849)	849		,860		,87			880		890		10F (0. 2 -	901
HOBHEGA	(819)	AACI	GTTCC	TACG	AAAGC	JAATC	TOTCI	TAA	AAGC	GTAP	GGAT	TTT	ACGC	CTC
BCV HE	(819)	AACT	GTTC	TACT	AAAGC	JAATC	Tere1	TAA	AAGC	GTAZ	GGAI	TIT	ACGC	CTC
MHV HE	(468)	AAGC	TTACC	CTCA	AAGGG	TATT	rocer	CCA	AAG	CAAT	KELC CC	1.1	A.L.C.	0.1
Consensus	(849)	AACT	GTTC	CTAC	AAAGC	CAATC	TGTCT	TAAT	CAAGO	GTAF	LGGAI			
													Section	
	(902)	902	Ş.	910		920		930		2	40			95
HOBHEGA	(872)	TAGA	GGTT	зттса	TTCG	CGGTG	GAACI	ATG	CAGO	CAG	CTG	TAA	CATG:	ACC
BCV HE	(872)	TAUA	GGTT	JTTGA	TICE	COCTO	GAAC	ATG	GAGG	BGAGI	CTG	TAYA	CATG.	ACC
MHV HE	(521)	TGCA	GGTA	STUGA	CTCAL	ACCTC	GAGT	GCA:	rock	CAGI	CAG	LCAA	TATG.	ACC
Consensus	(902)	TACA	GGTT	GTTGA	TTCG	CGGTG	GAACI	ATG	CAG	CAG	CTG	TAA		
													Section	n 19
	(955)	955	960		,970		,98			,990				100
HOBHEGA	(925)	GCGG	TTCC	TTGTC	AACC	rccgr	ACTG:	TAT	PTTC	TAA	PRC/P2	GTA	CCAA	CT.
BCV HE	(925)	GFAG	mmac.	ተጥር ጥር	AACC	CCCC	ACTO	PTAT	rrrc	TAA	PTCT7	ACTÁ	CCAA	TT:
MHV HE	(574)	GCTG	CAGC	CTGTC	AGCT	GCCAT	ATTG	PTTC	TTTC:	CAA	CACA	CTC	CGAA	TT.
Consensus	(955)	GC G	TTGC	TTGT	CAACC	CCGI	ACTG'	TAT	TTTC	GTAA!	PTCT	ACTA	CCAA	TT
													 Sectio 	n 20
	(1008	1008		.1	020		1030		,104			1050		106
HOBHEGA	(978	тотт	GGTG	TTT	-ATG	TTATA	AATC	Argd	AGAT	COTG	GTTT'	TACT	AGGA	TA
BCV HE	(978	TGTI	COTO	TTT	ATG	ATATO	AATO	ATCC	GGAT	CCTG	GTTP:	TACT	AGCA	TA.
		TAGI	CCTC	GCACA	ACATG	ATGCG	CACC	AFGG	TGAT	TTTC	ATTE	CAGG	CAGT	TA
MHV HE	(627				ATG	ATAT	AATC.	ATGG	GAT	GCTG	GTTT'	TACT	PAGCA	TA.
MHV HE		TGTT	GGTG	TTT										n 2
MHV HE) TGTI	GGTG	TTT									Section	n
MHV HE Consensus	(1008	1061		.1070		,1080)		090		1100			111
MHV HE Consensus HOBHEGA	(1008)) <u>1061</u>	TGGT	,1070 TTGT:	PATAT	AATT	ACCT	TGTT	TTTC	GCAG	CAAG	GCG'	eèrri	111 AG
MHV HE Consensus HOBHEGA BCV HE	(1008 (1061 (1028	1061 TTAC	TGGT	,1070 TTGT: TTGT:	PATAT	AATT GACT	ACCI	PGTT TGTT	TTTC	GCAG	CAAG CAAG	GTG	eêrtî erri	111 AG
MHV HE Consensus HOBHEGA	(1008 (1061 (1028	1061 TTAC	TGGT	,1070 TTGT: TTGT:	PATAT	AATT GACT	ACCI	PGTT TGTT	TTTC	GCAG	CAAG CAAG	GTG	eêrtî erri	111 AG

- Section 22									****
1166	1150		,1140		1130	<u> </u>	,1120	1774	(1114)
TOTOTOAC	TATOCCACA	TACCCC	GCCTCTC	TCTG	CAGTG	GTTAG	GATAAT	TAT (HOBHEGA (1081)
maccoom.	TATOCCACA	TACCOT	GCCTCTC	PCEG	CAGTG	GTTAC	GATAAT	TAT	BCV HE (1081)
TOTCCARC	TACGGICAT	TA CCC	GCC True	TCTG	CAGTG	GTTAG	GATAAT	TAT	Consensus (1114)
TGTCC AC	INIGGCAGA	INCCC							
Section 23		4000	1100		190	4		1167	(1167)
1219		,1200	1190		100	m A min a	Padmak	200	(1167) HOBHEGA (1134) BCV HE (1134)
TECGCTAC	TGTGTATGA	CATTTG	ATTTACC	CIG	ALAAC	LATIA	100154	700	BCV HE (1124)
TEEGCTAC	TGTGTATGA	TATATG	ATGTACO	OGTG:	ATACA	TATTA	TECTOA	200	BCV HE (1134)
ጥሮሮርሮሞልር	TGTGTATGA	TATTTG	ATGTACC	CCTG	ATAAC	TATTA	rgctga	TGC	Consensus (1167)
0									
4070	1260	1250		.1240		1230		1220	(1220) HOBHEGA (1187) BCV HE (1187)
12/2	1200	PHO COC	ringace	remin	GGGAT	TGCTT	LTATTT	CAG	HOBHEGA (1187)
THETAGIL		WA COMO	TTCCCC	TTTA	CETET	TOCTA	CATAC	CGG	MHV HE (836) Consensus (1220)
LEGEGTET	I G TU GAUUA	MMCCCC.	PTOCOMO	Commi	GGCAT	TGCTT	TATTT	CAG	Consensus (1220)
TTGTAGTT	TCATAATTA	TTGCGG	1166616						
- Section 25					400	0	,128	1273	(1273)
1325	1310		,1300) ((2)	,129		figigation 5	mma	HOBHEGA (1240)
GCTTAGAC	STGUATGAT	actacco	PARTICET	CGA	PATEG	a sevantar.	LIGITA	1,11,1	HOBHEGA (1240) BCV HE (1240) MHV HE (889)
SCTTAGAC	TIGCATGAT	ACTAGGO	PAATGGT	GGA:	TATE	LATT	LIGIIA	4.4.6	MHV HE (1240)
SCTTAGAC	CTGCATGAT	ACTAGG	PAATGGT.	PGGA:	TATGG	TATTT	TTGTTA'	TTG	Consensus (1273)
Section 26									
						1337	_	1326	(1326)
					SEQ II	AAC	ATCTA	CAT	HOBHEGA (1293)
			9885	NO:	SEQ II	AAG	ATCTA	CAT	BCV HE (1293)
			9896	NO:		35. T 15.25	· · · · · · ·		MHV HE (940)
							י אידים ייני אי		Consensus (1326)

FIGURE 4

FIGURE 4A

											- Secti	on 1
	(1)	1		1	0		20					3
avian infectious bronchitis poi 1ab	(1)	101714777	- cariro									
bovine coronavirus pol 1ab	(1)	MSKI	NKY	GEE	L HWA	PEF					SSSEV	
Human corona 229E pol 1ab	(1)						M2	CNI	VTLI	VAS	DSEIS	ANG
Murine hepatitis pol 1ab	(1)	MAKM	GKX	GIG							ERSEE	
Consensus	(1)	MAKI	ΚY	GL	WA	PEF	PWM	NZ	EKI	. NP	DSSE	
											- Section	on 2
	(40)				.50			0				_ 7
avian infectious bronchitis pol 1ab	(1)							- I-				
bovine coronavirus pol 1ab	(40)	CSTT	AQK	Æ T	GGIC	PEN	HVM	DCF	KLIE	QEC	cvoss	Βij
Human corona 229E pol 1ab	(21)	CETI	AQA	VRR	YSEA	Asp	GFR	CRE	vsiii	LQD	CIVGI	AD
Murine hepatitis pol 1ab	(40)	GPSA	ΑQE	PKV		LVN	HVR	NCS	REPA	LEC	CVOSA	TE:
Consensus	(40)	CST	AQ	LK	G	N	HVR	7 C	RLL	LEC	CVQSA	II
											Section	on 3
	(79)				90			100				11
avian infectious bronchitis poi 1ab	(1)											
bovine coronavirus pol 1ab	(79)	ETVM	NTR	PYD	EEVL	LQD	TEO:	ŒΙ.	VEV	PL.	GMSTE	AC
Human corona 229E pol 1ab	(60)	TYVM	GLH	GNQ	TLFC	NIM	KFSI	PP	MLHO	;	WL	V.F.
Murine hepatitis pol 1ab	(79)	DIFY	DED								RUSIQ	
Consensus	(79)	DIVM		PN	LE	IM	ALQ	R F	ATA	P	LSI	
	***			-							Secti	
and an internet and beautiful and a	(118)	118	07 h	754	130)		,14				15
avian Infectious bronchitis pol 1ab	(1)		MAS	SLK	OGVS	PK-	. 10713701	Les war		PRD	VIIVS	ΚŊ
bovine coronavirus pol 1ab	(118)	VRQC	NPN	GWI	MODE	RRR	SVC	IT G F	CAVI	KHV	AYOLY	MI
Human corona 229E poi 1ab Murine hepatitis poi 1ab	(93)	NSNY	LE	EMD	VVFG	NR-	-00054-0	- QC	GNY	YTD	OYLCG	AD
	(118)	Mrev	WIE K	EAA	MCHE	KRV	CECI	IJYR E	CSCI	AHV	AEHLE	TV
Consensus	(110)	N G	LР	SF	MGLE	KR	LCI	ITG	CAV	HV	AYLLF	
	(467)	457									Section	
avian Infectious bronchitis pol 1ab	(157)	10/	21727027			70		!	80		199	19
bovine coronavirus pol 1ab	(24)	REQU	EDA	LFF	YTSH	NPK	DYAI	ALI	VROF	EDR	SLOTE	KQ
Human corona 229E pol 1ab	(107)	PAUV	CLG	AGC	EN GW	A.T.F	DA E	TPV(SKKI	IVE	VMYI. WLTKR	$\mathbf{R}\mathbf{S}$
Murine hepatitis pol 1ab	(120)	KPVM	200	TMC	b VIDH	EGE	NEE	ILL	GHT	VCA	WLIKK	KP.
Consensus	(157)	PDGV	ពីកិច្ច	NGR	(II) T.G.M	EAR	V I A	BEZ	AKOV	HOP	WSILL	RK
Consensus	(101)	PDGV	CDG	Tre C	i F. v G W	FIP	L A.	LPIN	ARQE	.1 P		
	(400)	400				040		_			Secti	
avian infectious bronchitis pol 1ab	(196)					210	-	123_2	220	779 725		23
bovine coronavirus poi 1ab		KEET				V	C(UFI	PKG	DKI	TPGVP	ΑK
Human corona 229E pol 1ab	(196)	GEKG	-AY	NKC	HKRE	GFE	HVY	FKA	CE O'A'	DDA	HDEPK	GK.
Murine hepatitis pol 1ab	(100)	DYKR			-55	~-Q	N3	ΉV	EEIF	YVH.	GDÄLH	ТĻ
Consensus	(100)	GHAG	9 4 T	SGH	LKKA	A TEN	PAYAYI	JE N	E LAC	EEX	HLNEK	ĿΚ
Consensus	(180)	GFKG			KRA	. м	1 A X J	1 T	EDA	DPA	HDAPK	GK

								Secti	on 7
and an infantament of the state	(235)		240		250		260		27
avian infectious bronchitis pol 1ab	(88)	LKAT	SKLAD	LEDIE	GVSPI	ARKYR	ELLKTÃO	COWSLT	VE
bovine coronavirus pol 1ab	(234)	SKKA	YALIR	GYRGI	K B T T. YA	IDDVC	COVECT	NDOTER	7 100
Human corona 229E pol 1ab	(190)	NGSV.	PEWER	EVKTS	SKVVI	SDALD	KTYKVFO	SPVMT	NC
Murine hepatitis pol 1ab	(235)	SCKA	YABLK	GYRGI	KPTEF	LOOVE	CONTRACT	AVOTO	niviv
Consensus	(235)	SKKA	YALAK	GYRGY	KPILLY	DOYG	CLYTGAL	LA GLT	Y
								Secti	on 8
	(274)	274	280		290		300		04
avian infectious bronchitis pol 1ab	(127)	LDVR	AOTLD	EIFDE	TET I WIT	ON A	KIHVSSN	ANDDE	VIC.
bovine coronavirus pol 1ab	(273)	DKTL	OEMKA	LECT	SOFFE	Director Co	AWHVVRI	Company	VG
Human corona 229E pol 1ab	(229)	NALE	AFTERP	VETSE	LVOCTO	CTKG	WSVGDW	C DO CIG	Z.F.
Murine hepatitis pol 1ab	(274)	DEPL	SEME	1.0.0176	in Misting	TET ST	AWHYDRI	i GERGIO	
Consensus	(274)	DITL	AFTKD	LFPTE	ISD I	DA A	AWHVDRI	A A A	N. S.
						D V V.	AWHVDKI	Section	KTI
	(313)	313	320		.330		240	Secil	
avian infectious bronchitis pol 1ab	(166)	Marian.	TADM	c offer o	DOG	11100	340 ARIFOKA	d1050	35
bovine coronavirus pol 1ab	(312)	SASM	Tookin	O S MILE	MERCE	KOOL	ike eyn	WAIFE	NVI
Human corona 229E pol 1ab	(268)	W.F. S. II	L TOTAL	DOME	TEDICI	VACA	AGAGIK)	YADDS	III
Murine hepatitis poi 1ab	(313)	TOTAL OF	The State of	PGNVN	FGHAVI	THUO	AGAGERY	FCGMI	UKI
Consensus	(313)	TAT A TO	VRGUU	A.C.D.	THURVI	EDAA	VREPAHI	LAANA	LVI
Consensus	(515)	VIAT	VECVE	AANĞE	TEDLVI	GSVV	AREPIKI	LAA S	IV
	(352)	252	. 36	~				Section	
avian infectious bronchitis pol 1ab					370	3_05000	380		390
bovine coronavirus pol 1ab	(351)	DODE	T IN DITA	CORREA	KCARSI	TAAA	ERTLYV	KEFAG	TCI
Human corona 229E pol 1ab	(307)	AND MEN	PMDT I	SCERM	HURY	NAFIX	SVDEKD	DEVMO	EGY
Murine hepatitis pol 1ab	(352)	STO.	- FW		EGV SVW	RVIA	QSVDCE	VASST	EVE
Consensus	(352)	RLPN	IIL	EIMPX	TUSSAT	EFCX	KTKLCLC	GELTQ	E G
Conscisus	(JUE)	KTEN	IIP	F. T					FGY
							KL DC		
	(204)	204					L KL DC	- Section	11
avian infectious bronchitic col 4ch	(391)	391	2.52.	400	4	10		Section	11
avian infectious bronchitis pol 1ab	(244)	ASIN	BAVAR	400 for a sti	A PNG FMG	10 SKT P	MATAIR PV	- Section	11 429
bovine coronavirus pol 1ab	(244) (390)	ASING	SAVAR ODLEB	400 DFEED	A PNGFMG BGMMTT	10 SKIF	TLAFFK	— Section	11 429 VVE
bovine coronavirus pol 1ab Human corona 229E pol 1ab	(244) (390) (336)	ASING	SAVAK QDLED	400 DFEET FKGWV	A PNGFMG PGNMID	10 SKIF	TTMFFK	— Section BAAVE EDGDL	11 429 VVE
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab	(244) (390) (336) (388)	ASING IBCE EEHVI VDCC	SAVAR DDLEB NRMDT SDTCD	400 LFEEL KGWV PCFNV	4 PNGFNG PGNMIT RNSVII AGNMMÉ	10 SKIF SEAC ECRL	TTAFFK TOCOHVY AMBOAEM	— Section EARVR EEGDL ITSNVR	429 VVE BAC RQV
bovine coronavirus pol 1ab	(244) (390) (336) (388)	ASING IBCE EEHVI VDCC	SAVAR DDLEB NRMDT SDTCD	400 LFEEL KGWV PCFNV	A PNGFMG PGNMID	10 SKIF SEAC ECRL	TTAFFK TOCOHVY AMBOAEM	Section EAAVR EEGDL ITSNVR MPWEL ESAVR	11 429 VVE NAC RQV EAC
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab	(244) (390) (336) (388) (391)	ASING TOCE TECHVI VOCC IDC	SAVAR DDLEB NRMDT SDTCD	400 FEED KGWV CFNV RGWV FKGWV	4 PNGFNG PGNMIT RNSVII AGNMMÉ	10 SKIF SEAC ECRL	TTAFFK TOCOHVY AMBOAEM	— Section EARVR EEGDL ITSNVR	11 429 VVE NAC RQV EAC
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab Consensus	(244) (390) (336) (388) (391)	ASING IPCE EEHVI VDCC IDC (SAVAK ODLEB NRMDT SDTEB GDLCD	400 LFEET KGWV L'CFNV RGWV FKGWV	A PNGFMG PGNMTT RMSVTI RONMME	SKIF SEAC SEAC SEPC	ette a Eggtknå Mogyen Eggtknå Eggtknå	Section EARYR EEGOL ITSNYR MPWEL ESAVR: Section	HACK LACK
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab Consensus	(244) (390) (336) (388) (391) (430) (283)	ASING EEHVI VOCCI IDC (SAVAR ODLED NRMOT SDTCD SDLCD	400 LFEET KGWV CFNY RGWV FKGWV	A PLIGENG PGMMIT RMSVTL AGNMME PNNMMD	10 SKIF SEAC ECRLA	TAFFK TCCHVY MEGAEM PGCTKNY PTLG Y	Section EAAVR EEGOL ITSNVR MPWEL ESAVR Section	11 429 VVE RQV EAC LAC 112 468
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab Consensus avian infectious bronchitis pol 1ab bovine coronavirus pol 1ab	(244) (390) (336) (388) (391) (430) (283) (429)	ASING FBCE EEHVI VDCC IDC 430 NIPNA SSGVI	SAVAR ODLEB NRMDT SDTOD GDLCD APRET	400 LFEE KGWV CFNV RGWV FKGWV A40 KGFEV	A PLIGFNG PGMMIT RMSVII AGNMME PNNMMD VGNAKE SAAGYG	10 SKIF GEAD ECRLA ECRLA ECRLA ECRLA ECRLA ECRLA A50 TQVVI	TAFFK TCCHVY MEGAEM PGCTKNY PTLG Y	Section EAAVR EEGOL ITSNVR MPWEL ESAVR Section	11 429 VVE RQV EAC LAC 112 468
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab Consensus avian infectious bronchitis pol 1ab bovine coronavirus pol 1ab Human corona 229E pol 1ab	(244) (390) (336) (388) (391) (430) (283) (429) (375)	ASING TBCE(EEHVE VBCC/ IDC (430 NIPN/ SSGVI	SAVAK ODLED NRMDT SDTOD GDLCD APRST CPVNP	400 TFBET FKGWV FKGWV A40 KGEEV	A PING FMG PGMMIT RMS VIL AG NMME PNNMMD VGNAKG SAAG YG YDDIFA	10 SKIF SEAC ECRLA SEPC OF C 450 TQVVI	TAFFK TCCHVY AMEGAEM FGCTKN FTLG Y VRGMEND	Section EAAVR ETGDL TISNUT MPWEL ESAVR Section LTLEDG	VVE VVE RQV EAC LAC LAC LAC VFC
bovine coronavirus pol 1ab Human corona 229E pol 1ab Murine hepatitis pol 1ab Consensus avian infectious bronchitis pol 1ab bovine coronavirus pol 1ab	(244) (390) (336) (388) (391) (430) (283) (429) (375) (427)	ASING TBCE(EEHVI VBCC(IDC (430 NIPN/ SSGVI ASGVI SSGVI	SAVAR ODLEB NRMDT SDTOD GDLCD APRET	400 FREE KGWV CFNW RGWV FKGWV 440 KGFEV VUHUK GWFDW	A PLIGENG PGMMIT RMSVTL AGNMME PNNMMD	10 SKIF SEAC ECRLA EEPC GF C' 450 TQVVV GFGC' E	TAFFK TCCHVY AMEGAEM FGCTKN FTLG Y VRGMEND	Section EANTR ETEGOL ITSNUT MPWEL ESAVR: Section LTLEDO GOTVV:	VVE VVE RQV RQV LAC LAC LAC VFC

								Section 1	
	(469)			480		90			0
avian infectious bronchitis pol 1ab	(322)	DIPVEP	EGWSA	LLDGH	CYVF	SGDRF	YAAPL	SGNFAI	ġ.
bovine coronavirus pol 1ab	(468)	GCVYWS EDFFGP	-PARN	IWIPI	LKSSVI	SYDCL	VXTGV	VÇCKAJ	Ē?
Human corona 229E pol 1ab	(405)	BDIFGP	-CWSA	LASAL	KOLKVI	IGELV	REVKS	ICNSAV	Ĥ
Murine hepatitis pol 1ab	(462)	SAVYWS	-PCPG	MWLPV	IWSSVI	SYSCL	TYTCV	VCCKAI	ĹΑ
Consensus	(469)	D VYWS	PWSA	INIPI	L SSVE	SYDGL	YTGV	VGNKAI	Ľ,
								Section 1	4
	(508)			520		530		5	
avian infectious bronchitis pol 1ab	(361)	DWHCCE	RVVCL	SUGMT	PEINDO	LILAA	LYSSE	SVSEL	7
bovine coronavirus pol 1ab	(506)	KEUNLO	CKALY	DAKOL	HKCGNI	HOREL	LGVSB	VWHEQI	en Light
Human corona 229E pol 1ab		VVGGTI							
Murine hepatitis pol 1ab	(500)	QETDA	CRSLY	MDYVO	HEGGN	EORAL	LGLDD	MYHROI	ď
Consensus	(508)							VWHEQI	
								Section 1	ıŧ
,	(547)	547		560		570		5	58
avian infectious bronchitis pol 1ab	(400)	ALKKGE	PFKEL	GHKFV	YAKDA	VSFTI	AKAAT	IADVL	Ŕ
bovine coronavirus pol 1ab	(545)	LNRGV	KPLLE	NIDYE	NMRRAI	KEGLET	ETVEA	DOFME	F
Human corona 229E pol 1ab	(482)	TAGKAI	DKVFE	YVLLD	NALVK	WITKI	KGVRE	ROLNK	ý
Murine hepatitis pol 1ab	(539)	VNRGD	SLIDE	NVDLE	VKRRAI	FACK-	PATES	DGLVPI	Ċ
Consensus		INRKA						DGLVPI	
							<u> </u>	Section 1	16
	(586)	586		,600		,610			62
avian infectious bronchitis pol 1ab	(439)	FOSAR	IAEDV	WSOFT	EKSFE	FWKLAY	GKVRN	LEEFM	K
bovine coronavirus pol 1ab	(584)	CODLY	RAYYI	AVEGO	AFCDY	ADMICE	AVVSK	KELL	õ
Human corona 229E pol 1ab	(521)	YATVV	GSTEF	VKSS-]	RVERSI	AVETI	ANNYS	Ķ
Murine hepatitis pol 1ab	(577)	DOGDVI	RSYVI	IKSĞÖ	AFTSM	MVNFSF	EMIDM	CMDMA	Ĺ
Consensus		LDSLVV						A EMS	K
								Section 1	1
	(625)	625	30		40	650			6(
avian Infectious bronchitis pol 1ab	(478)	YVCKA	MSIVI	LAAVL	GEDIM	HÚV3	- QVIIIY	KLGVL	Ê
bovine coronavirus pol 1ab	(623)	SLDSL	SAATH	LNSKI	VDLAO	HESDEC	TSFVS	KIVHE	É
Human corona 229E pol 1ab	(554)	FDEGY	VVIGI	VAYEN	SDGYE	RUMASI	NSVLI	TAVYK	P
Murine hepatitis pol 1ab	(616)	EMHDV	VATK	VKKYT	GKLAV	REKAL	VAVVE	KTTEW	Ĩ
Consensus	(625)	FLD L	LAVE	LAAVI	GDLAF	RLMA C	SVVS	KIVHE	Ē
								- Section *	1
	(664)	664	670		680		90		7
avian infectious bronchitis poi 1ab		KVVDF		GPCVO	EKRAK	LIVTÊ	FCVLE	GVAOH	c
bovine coronavirus pol 1ab	(662)	TFTTS	PALAEZ	AWVLEH	VIHGA	YIVVE	DIYEV	KNIER	5
Human corona 229E pol 1ab		FAFN	NVMGTÍ	PEKEP	TTVTC	ENESS	VLEVI	DKLTE	Ì
Murine hepatitis pol 1ab		LAVEL							

												S	ection	19
	(703)	703	19762	710	2000000			0			730			74
avian infectious bronchitis pol 1ab	(553)	QLL	LDA	CHSI	YKS	FK	KCA:	LGR	THO	DLL	FWK C	GVI	IKIV	QI
bovine coronavirus pol 1ab	(701)	SAV	AOAI	RSV	AKV	VL	DSL	RVT	FIL	GLS	CFKI	GRI	RET	T.
Human corona 229E pol 1ab	(701) (632)	T DX	SID	/IDN	EII	VK	PNI.	SLC	VPI	YVR	DYVI	KWI	DEC	R
Murine hepatitis pol 1ab	(054)	LNE	AHVI	A.A.	TEV	$(\pm i)$	DSM:	5.V.S	LED S	GLT	VVKI	ASI	IRV.	Li
Consensus	(703)	v	SDAR	KSL	FKV	VK	DSI	SVS	II	GLS	FK	Ğ	RIC	
						-,						Se	ection	20
	(742)	742		.750)		- 2	760			.770			78
avian infectious bronchitis pol 1ab		GD-												_
bovine coronavirus pol 1ab	(740)	CSK	IYEN	ERG	LIGH	SS	OLE:	Dv	YDI	FM P	sove	KAI	OKE	т,
Human corona 229E pol 1ab	(6/1)	Y37-							- 0222	25,				
Murine hepatitis pol 1ab	(733)	GSK	YYE	EVOK	STIS	AY	VMP	ZGC	SEP	MCT.	VGET	ED.		
Consensus	(742)	GSK	IYE	ĵ –	L		LP		D					
												_ Se	ection	21
	(781)	781		7	90			800	1					~ 4
avian Infectious bronchitis poi 1ab	(594)				P Tie	FD	ATO	200	SEP P	TOTAL	VOER	S To		72
povine coronavirus poi 1ab	(779)	LKG	SGST	TEST	ABC	TAXE	EX. 27.233	rim e	Transition to	ישויים	CONTRACT.	Car.	TO TE Y	
numan corona 229E por 1ab	(6/3)			N	R S D	THE RE	mnvi	ים וליכ	THE TAI	retended	THE PART OF	27.75	in ma	**
Murine hepatitis poi 1ab	(766)			AVE	EDE	VV	DVV	A P	7 T V	OCC	CKDI	me :	E V I	
Consensus	(781)				EDN	VV	DVV!	A	T.T	LGT	SDPP	CT7	DV.	330
										201	BULL		ection	
•	(820)	820			830			8	40					85
avian infectious bronchitis pol 1ab	(623)	DVT	EPEN	IOPG	ний	OI	EDD	KN	VMF	FRE	KKDE	NTI	VIDE	M
bovine coronavirus pol 1ab	(818)	VDN	v YMZ	KAC	DKY	ΥP	VVV	7G-	HAVE	13716	O A TOP	770	经常	SAR.
Human corona 229E pol 1ab	(703)	FWD	TIVE	POP	STI	KV	TDG	Se T	DIN	TOWN Y	NITTATO	MOI	MIT	
Murine hepatitis pol 1ab	(798)	V DK	YMA	Rec	DOE	N 13	WWX	ON U	TXTO	X7 1	CHE	PIOR	10 THE	2
Consensus	(820)	VV	LYMA	KCG	DTV	YP	VVV	SK.	WVC	VID	O M D	VDC	MAN	E.
	,,								****	V 11 D	2 111	.VEC	ction	23
	(859)	859			870	1			880					89
avian infectious bronchitis pol 1ab	(662)	OTTG	TNY	TATO SE	AZC	K T	roler	- D.M.	milio	OT D	D D O T	TID.		09
bovine coronavirus pol 1ab	(856)	TEK	OPT	MNE	TAS	T P	C marie	WE	VIE T	PKD	PNTT	NO	NOC	4
Human corona 229E pol 1ab	(742)	KEN	LTOC	GLL	GTC	AK	RFKI	WT.	CET	T. F A	V LID E	THE DA	PUAT O	W.
Murine hepatitis pol 1ab	(837)	EFW	DKPK	ZRK	CPS	efe-a	REST	a m	FAI	ייי מ'כו	en ev	200	AC G	- 6
Consensus	(859)	LN	D P	VK	TAS	T	R T T T	TT	FIL		FNSV			
											LNSV		ction	
	(898)	898			٥	10			.920	`				กวง
avian infectious bronchitis pol 1ab	(701)	CCC	PPWN	ı – – –		_ mr	DEKT	Chv	<u> </u>	T P 12	DATE OF	mir	- K-	23
bovine coronavirus pol 1ab	(895)	nun	TY T	MER	EVA	17.47	10.00	IN P	ハムド ン部等	TEV	2101	COMP	Ship.	4
Human corona 229E pol 1ab	(895) (781)	KU C	33 + 14/1 T		- IM	PANS	かまがた	12 12	400	FOR	كالطبدوة	V.G.	IK VIS	γVI
Murine hepatitis pol 1ab	(875)	EVD	K DŠŽ	The second	T. T. D	77Y.7	66	EL	H T	E DIG	L'XIT A	THE	TAC	K١
Consensus	(898)	EVC	D Λ.	T.DP	טטט	VV	E D V .	CE	Tri S	L'GW	EHDV	1,5,1	AVC	A1
22.10011040	,,	_, _		200		* V .	·		T 77.2	FUK	⊾nu∨	107	KVC	AI

									_ Section	n 25
(93	37)	937			,950		,960			975
avian infectious bronchitis pol 1ab (73	33)	LOKLET LOKLET ENKTE!	DD	LK	PEAPE	PPPF	ENVAL	VDKNG	KELDO	IKS
bovine coronavirus pol 1ab (93	34)	LOKLET	ONS	LE 5	DEAGE	EVLA	SKLYC	AFTAP	EDDDF	LEE
Human corona 229E pol 1ab (80	07)	ENKTE	EW	TERR	PHNDE	IKSF	STFES	AYMPI	ADPTH	FDI
Murine hepatitis pol 1ab (91	14)	DORLAC	DY	VYTH	DEGGE	EVIA	PRMYC	SESAP	DDEDC	VAA
Consensus (93	37)	LNKLEA	DW	LFLI	FPEAGE	EVIF	SKLYC	AFSAP	DDDDC	IDA
									- Section	n 26
(97	76)	976			,990		.100			1014
avian infectious bronchitis pol 1ab (77	72)	CHILLA	ŧĎΥ	ESD	DIEEE	DAEE	CDTDS	GEALE	CDTN-	
bovine coronavirus pol 1ab (97	73)	SGVEEL	ΧDV	EGE	TOLIN	TSAG	PCVA	SEOEE		
Human corona 229E pol 1ab (84	46)	EEVELI	ĎΑ	EFVI	PGCG	PEAV	ĨĎЕнŸ	FYKKD	ĞVYYP	
Murine hepatitis pol 1ab (95	53)	sgveei Eeveli Dvvdai	žΕΝ	O D	AEDSZ	VIVA	DTOEE	DGVAK	GOVEA	DSE
Consensus (97	76)	D VE I	ดีดี	E DI	DDSA	ILAA	DD DA	E EE	G	
					<u> </u>				- Section	n 27
(101	15)	1015 1	020		.10	30	.10	040		1053
avian infectious bronchitis pol 1ab (80	07)			SECI	EECDEL	TKVL	ALTOD	PASTK	YPLPL	RED
bovine coronavirus poi 1ab (100	04)	S	ĔΙ	LED!	T L D DG E	CVET	SDSÖV	EEDVE	MSDFA	DLE
Human corona 229E not 1ab (88	R21		ũn α	TOTAL TO	PVART	KAAC	CRAZER	SHOWE	TERRE	DITTY
Murine hepatitis pol 1ab (99	92)	ICVAH	GS	OBIO	CAPPOZ	VGSQ	TPLAS	ABETE	VGEAS	DRE
Consensus (10°	15)	Š	รั	SEDI	L EDD	AA	A IQ	AEDVE	V D A	DLE
									_ Section	n 28
(105 avian infectious bronchitis poi 1ab (83	54)	1054	,106	0	.1	070		1080		1092
avian infectious bronchitis poi 1ab (83	38)	YSVYNO	CI	KHK	DALDAN	NLPS	Ğ			
having accompained and date (400	201	-	4.6	2723_0	A5	-3457				
Human corona 229E pol 1ab (9	16)	RVKLC	ÉF	EDE	KLVDVI	EKAT	ğ			
wunne nepatitis poi Tab (To.	31)	GEALA	CAT	V CA3	DAVEDAL	PDOV	EAFEL	EKVED	SILDE	LQT
Consensus (198	54)	VI I	FE	vc i	DAVDVC	P I	G			-
									_ Section	
avian infectious bronchitis pol 1ab (86 bovine coronavirus pol 1ab (105	93)	1093	.1	100		,1110		1120		1131
avian infectious bronchitis pol 1ab (86	60)									-EE
bovine coronavirus pol 1ab (105	57)		-						EF	VKV
Human corona 229E pol 1ab (9)	38)								K	KIK
Murine hepatitis pol 1ab (10)	70j	ELNAPA	ADK	TYE	DVLAF	DAVCS	EALSA	FYAVP	SDETH	FKV
Consensus (10)	93)									ĸv
									- Sectio	
avian infectious bronchitis pol 1ab (8	32)	1132		1140		,1150		,1160		1170
avian infectious bronchitis pol 1ab (86	62 <u>)</u>	TFVVN-					-NCFE	GAVEP	EPOKV	V.D.V.
bovine coronavirus pol 1ab (10	62)	LDLYV	EKA	TŘN	NCWER	VEAV	MOKER	COFKD	KNLOD	LWV
bovine coronavirus pol 1ab (10) Human corona 229E pol 1ab (9)	42)	HEG			-DWDSI	CKTI	QSALS	VVSCY	VNLPT	YYI
Murine hepatitis pol 1ab (11	09)	CGFYS	ĔΑΙ	ERT	NCWLR	STRIN	MOSLP	LEFKD	LEMOK	LUL
Consensus (11	32)	DLX	P	R	NCWLRS	LV	MQALP	L FKD	LNLQ	LWV
'•										

						Section 31
(1171)	1171	,118	0	,1190		1209
avian infectious bronchitis pol 1ab (884)	EGDWGI	BAVDAQ	EQLCQQ	EPHOHTF	EEPVENS	TGSSKTMT
bovine coronavirus pol 1ab (104) Human corona 229E pol 1ab (972) Murine hepatitis pol 1ab (1148)	T-XKQQ	YSOLF V	DTLVNK	TLANEVV	POGGYVA	DEAYWELT
Human corona 229E pol 1ab (972)	YDEEGO	SNDESE	PVMISE	WEDSVOO	ACCEATL	PDIAEDVV
Murine hepatitis pol 1ab (1148)	SYKAG	POCTV	DKLVKS	VEKSIIL	POGGYVA	DEAYEFES
Consensus (1171)	LYK G	YAQLEV	D LVN	IPLSIIL	PQGGYVA	DFAYFFLT
						- Section 32
(1210)	1210 '	,12	220	,1230		1248
avian infectious bronchitis pol 1ab (923)	EOVVVI	EDGELP	VIVE ON C	DVVVYTP	TDEEVAK	ETAEEVDE
Dovine coronavirus pol 1ab (1140)	LCDWQC	CVAYWK	CIKCLI	ALKEKSI	DAMEFYC	DOVSHVCK
bovine coronavirus pol 1ab (1140) Human corona 229E pol 1ab (1011)	DOVEE	VNSIDD	IETV	KHDWSPF	EMPEEEL	NGLKILKO
wurine nepatitis poi 1ab (1187)	OCSIFIKA	AYANWR	CMT SEZONA Y M	FEKTOGE	DAMERYC	CHARLET CHARLES
Consensus (1210)	DQVF	A WK	CIECDL	DLKL GL	DAMFFYG	DVVSHVCK
						- Section 33
(1249)	1249		1260	1270		1287
avian infectious bronchitis pol 1ab (962)	FILIFA	AVPKEE	VVSQKD	SAQIKQE	PIQVVKP	QRE-KK
Munite nepautis por Tab (1226)	LANEM:	PERSAD	I P Yere P	FGVRDDK	HC A BWMP	REVERDANCE
Consensus (1249)	CGNSM	TISAD.	VPFTLH	GALKDD	FCQFVTP	RKVFKAAC
						Section 34
(1288)	1288		1300	,131	0	1326
avian infectious bronchitis pol 1ab (998)		ra emergranian	Charleston of the			XK
bovine coronavirus pol 1ab (1218) Human corona 229E pol 1ab (1082)	ANDANI	SHSMA	AADOKO	EDDHRVI	SITSDE	DEDIGHEM
Musica baselila and 410 (1082)		the contract management				
Murine hepatitis pol 1ab (1265)	AVDVNI	CHEMA	VVEGKO	LDGKVVT	KFIGDEF	DEMVGYGM
Consensus (1288)	ADAMI	HSMA	VVDGKQ	ID VT	DKF	DFIIGHGM
						- Section 35
(1327)	1327		1340		350	1365
avian infectious bronchitis pol 1ab (1000)	KEKVKI	ATCEK	PKFLEY	KTCVGDL	TVVEAKA	DEFKEFC
bovine coronavirus pol 1ab (1257)	SESMIT	FELAO.	LYCSCI	TPNVCFV	KGDIIKV	SKRVKAHV
iviurine nepatitis poi 1ab (1304)	TESMSI	PELACI	LYGSCI	TPNVCEV	KGDVTKV	VRIVMATO
Consensus (1327)	SFSMSE	PFEIAQ	LYGSCY	TPNVCFV	KGDIIKV	LKLVKAEV
						Section 36
(1366)	1366		1380		1390	1404
avian infectious bronchilis pol 1ab (1039) bovine coronavirus pol 1ab (1296) Human corona 2295 pol 1ab (1114)	TVNAAN	EHETH	GSGVAK	ATADEC	LDEVEYC	EDYVKKHG
bovine coronavirus poi 1ab (1296)	VVNPAN	IGHMAH	EGGVAK	ATAVAAG	COLAKEL	TDMVKSKU
Munne nepatitis pol 1ab (1343)	IVNPAN	IGREAH	3 AGVAG	ATAFKAC	SAFTKER	SOMURACE
Consensus (1366)	IVNPAN	IGHMAH(GAGVAK	AIAE AG	A FVKET	TDMVKAHC

							Section	n 37
(1405)		1410		1420	1	430		144
avian infectious bronchitis pol 1ab (1078)								
bovine coronavirus pol 1ab (1335)	VOAT	GDCYV	TGGK	LCKTVE	VVGPD	ARTOG	KOSYA	Th.
Human corona 229E pol 1ab (1153)	TCLN	CNAPRI	CTIR	OLOGIT	I E KOOK	PF.P		VN
Murine hepatitis pol 1ab (1382)	VCOV	GECYE	AGGK	LCKKVL	NIVEPT	ARGHG	KOCYS	T. F.
Consensus (1405)	VCQ	GDCY S	GGK	LCK VL	NIVGPD	AR G	KO YA	
							- Section	
(1444)	1444	.1450		.1460		1470		148
avian infectious bronchitis pol 1ab (1082)			-LVT	PSPVK	TOCUNN	VMCPP	HCDNN	Tu
bovine coronavirus pol 1ab (1374)	RVVK	HENKY	CVVT	THUSAS	rrevae	BUSTO	VIII CI	AE
Human corona 229E pol 1ab (1185)	VSEV	VKPVO	STER	GAMSCE	HYOTNI	YSONL	CVDGE	CV
Murine hepatitis pol 1ab (1421)	BAYO	HTNECT	SNIVER	OTTODA	E COL DO	DUGAM	V PT CA	r True
Consensus (1444	RAY	HINKCI	TVVT	TT. TSAG	TESVES	DUSLT	VILLE	T.
						2,021	_ Section	
(1483)	1483	1490	,	1500		.1510		152
avian infectious bronchitis pol 1ab (1110	KTVA				THE T. PHY		ZX E TIT	
bovine coronavirus poi 1ab (1413)	OVU	USWNOT	Speni	SKOOT	faveco	RKTAR	DISPA	ixic
bovine coronavirus pol 1ab (1413 Human corona 229E pol 1ab (1224	RTOP	DITA	Name of	DATATE	CIKDAD	VNAKS	PICTO	PT
Murine hepatitis pol 1ab (1460)	NAME	USNNOT	POPOV	FERCON	BEWARD TO	2 A	OTAK	TC
Consensus (1483	KVVL	VSNNOI	DFDV	TAKCOT	TIVDGT	KALAL	KLSIN	IT.T
							- Section	
(1522)	1522	.15	30	.154	0	.1550		156
avian infectious bronchitis pol 1ab (1149)	FAFE	CCTTRI	JATES					
bovine coronavirus poi 1ab (1452)	STORY	E TEDEN	CHARTIE	NEGATIV	STENT	OF UTS	THUN	AT
Human corona 229E pol 1ab (1256)	NTVD	TTPKET	FWVK	EKENAF	LVEDNA	AFYOG	DVDT	VN
Human corona 229E pol 1ab (1256 Murine hepatitis pol 1ab (1499	DVKF	VINAC	SIES	-ESCEV	SSYDVI	OFVEA	CRHD	Ot
Consensus (1522	DIVE	TA	LLFS	DL FV	SSHDVI	ODV A	LRHDI	i I.
							- Section	
(1561	1561		1570	.15	580			159
avian infectious bronchitis pol 1ab (1163								
bovine coronavirus pol 1ab (1491	DDAR	TEVOS	W.Fiv.W	SE CORN	VNEDYC	TNEWR	TWKY	r C
Human corona 229E pol 1ab (1295	VOED	FIVNA	NENE	AHGGGT	AKALDV	YTKGK	LORILS	KE
Murine hepatitis pol 1ab (1537	DDAR	VEVOA	MOCE	BTDWRL	VNKEDS	VDDVR	TKY	EC
Consensus (1561	DDAR	FVOAL	MD L	P GWRL	VNKFD	I GVR	TIKYI	FEC
							- Section	
(1600	1600		.1610		1620			163
avian infectious bronchitis pol 1ab (1163		OEHTÖ				FDV		
bovine coronavirus pol 1ab (1530	COND	fcson.	ZVEZY	VÖÖGSÉ	NKAPAT	BIRKE	FYOR	žĎŤ
Human corona 229E pol 1ab (1334	TGIA	G	w.anaca	KV	KVGTGV	MARCI	STRT	N
Murine hepatitis poi 1ab (1576								zbv

Section 43							· · · · · · · · · · · · · · · · · · ·
1677	,1660		,1650	- A	1639	(1639)	
OVYAKN	QFG	PGDSLG	SIVLKI	EVKYP	LTE	iab (1182)	avian infectious bronchitis pol
ECDGVNVTKHKCD	CONTRACTOR OF THE PARTY OF THE	TOTAL CALL	M D PARDY	TO MAKE T	33/1152	lab (1569)	bovine coronavirus nol
FONCTNING	KUTHRUFC	EXEVE	SICCIVAF	GVNFR	CEV	1ab (1615)	Munne nepatitis poi
FCDGINVTKHKC	KSLGSVFC	GESFG	SILVK	GVNFR	LTV	sus (1639)	Consen
Section 44							
	.1700	0	1690		1678	(1678)	
VARCOUNT OFFICE TIES	PRETTPUM	. VITDTT	FDKERT	TADDV	KIV	lab (1211)	avian infectious bronchitis poi
AT EXECUTE TO THE RESERVE	TO DO GO CATES	CORDIN	OFDINTE	CKVER	TINY	lab (1608)	povine coronavirus poi
UCKUKULUEGIIN	TIVEDERVE	C HI WELLER W	DVCNTK	T. F. M. I. I.	EET	lab (1399)	Human corona 229E pol
GEDEPOLUKYYTM	THE WALL THE THE	S END TOTAL	OMSDE	CKVPE	A	ab (1654)	Murine hepatitis pol
FDLKDLLAYY M	AVSDSE E	SEVDT.	OF NIS	GKVFF	IIY	sus (1678)	Consen
Section 45			2				
4700	.1740	30	.173		1717	(1717)	
Mo object of the second	TO DOMESTICA	THE POT TOTAL	OVEDNICE	OKENI	OPL	lab (1250)	avian infectious bronchitis pol
TO SHIATIONAKI	A WATER CHIEFE	PPRE	VENCK	SKWOV	EVN	ab (1647)	bovine coronavirus poi Human corona 229E poi
MAKA 2 OF METOS TIME	MANAGEN	CALLEY	F PM_ PV	EOPKIT	VOK	lab (1438)	Human corona 229E pol
INVACUMLONLSL	LO MENTAL OF THE PARTY OF THE P	FAFE	MVCGW	CKNSV	TG-	ab (1693)	Murine hepatitis pol
INVACLMLQAL L	MANAGETAL	A E I E R U	VERCHY	KWNV	T.	sus (1717)	Consen
Section 46	SWMMMCETM	LLIENQ	· LIGHT				
0 470	1780	1770	1		1756	(1756)	
USC CON A WATER STOR	DETERMINATE	- Werdish	WAKTTG	FIT. TEA	REK	lab (1289)	avian infectious bronchitis pol
INSCIANVISEESE	DEVENO I	of Charle	ATT DUD	Vallag		lab (1686)	bovine coronavirus pol
ATTELY OF POTTY POTTY OF THE	TO CHANGE IN ATTICAT	4T TO T TO	TEPDOM	DEPTU	CVA	(an 11476)	numan corona 229E noi
Lakosekeneesd	DEVENTER	SESSEDIT.	NUMBER	MOMOR	KEP	ab (1731)	Murine hepatitis pol
LAKGGFKFGDPSD	PRINCE	D C C K D	AM FED	FOWOR	KFK	sus (1756)	Consen
Section 47	ME VALVUAL	KBGKE		- 21121			
200	.1820	.1810		1800	1795	(1795)	•
ALCO THE	TE PER DATO MELL	VERN P	CHE TO TO T	TO MITTER	OME	ah (1327)	avian infectious bronchitis pol
BRCGIKS	THE REAL PROPERTY.	CARTED N	SOVETE	TERRITOR	SRD	ab (1725)	bovine coronavirus pol
TOWN TO THE WOR	CHARLEN	TABLEDIC	- CALMITAT	NKATO	CVD	ab (1515)	Human corona 229E pol
JGVKQEQRKGVDA	TO THE THE PARTY OF THE PARTY O	E CONTRACTOR	PEANTS	MRXIVIT.	STD	ab (1770)	Murine hepatitis pol
CGVKQEQRKGVDA	TRIVENCE	SENTEN	TOTE	T. P. V.V	STD	sus (1795)	Consen
Section 48	DELACKCE	JGA C	HD113	21(1)	010.		
	.1860	.1850		.1840	1834	(1834)	
1860 1872	1861,	7 TO DITE	##AC	T.P.C.		ab (1360)	avian infectious bronchitis pol
FRIGISNOPTOGA	CATNLLHER	TUPVR	DATE	DAG	17 M 17	ab (1764)	bovine coronavirus pol
THOUREDVEETTO	SUGKKLIHO	2 1 I ADG	والاطرارات	ODOR	DAIN	ab (1754)	bovine coronavirus pol Human corona 229E pol
TTETATESPECNA	TITPADYILI	TCDLVG	K P IN KAPAN	QKC1R	DND	ab (1800)	Murine hensilie nol
VHCTOPNURFLIC THCL FNUPTLIC	TEGSKLVHO	XNEAC	PRTAKE	CTT DK	VALID	an (1009)	warne nepanis por

							 Section 49
) 1873	1880	100	,1890		,1900	191
vian Infectious bronchitis pol 1ab (1393) · NNT DE	VIEASI	PYIL	LFATDO	PATV	DCDEDAY	VGTVV FV
bovine coronavirus pol 1ab (1803	SNTEA	SYRLER	GVGS	ANDEKE	DKVG	нүүникс	CEQSYQL
Human corona 229E pol 1ab (1590) SFVGE	LKAAEA	KVIT	IKVIEL	GVNV	HDVTVT	FDKSFEQ0
Murine hepatitis pol 1ab (1848) SNTPE	GRKUPE	DVVA	ANIFTC	GSVG	HYTHVKO	KPKYOL
Consensus (1873) SNTPE	LRKLP	VIS	ANITDO	GAVV	HYVHVK	CD SYQL
							Section 50
(1912	1912	,1920		,1930		.1940	195
vian infectious bronchitis pol 1ab (1432) STNSG	HCYPO	ACQA	FONLAP	DRKE	GKKSPY	TAMY.TR
bovine coronavirus pol 1ab (1842	DASNA	KKVTDV	TANI	SPOLYI	KNIK	OFFKSV	CTTYYID
bovine coronavirus pol 1ab (1842 Human corona 229E pol 1ab (1629 Murine hepatitis pol 1ab (1887) VEVIA	DKDKDI	SEAV	PSDTINT	SEEL	TKAIDW	DWVEFYG
Murine hepatitis pol 1ab (1887) DACNA	NKVSET	KONE	EDCLY	KNLK	OUESSV	LTTEXED
Consensus (1912	DA NV	KVTD	SGNI	SDCLYI	KNLK	QTFSSV	LTTFYLD
							- Section 51
(1951) 1951	,19		,19			198
vian infectious bronchitis pol 1ab (1471) AFKNE	TSLPM	KOSE	GKSKSV	KEDV	SNLATS	SKASFÓN
bovine coronavirus poi 1ab (1881) VKKI	YN PDE	OYYC	DECKY	CORT	TKAOPK	TEEKVDG
Human corona 229E pol 1ab (1668) KDAVI	TATVOI	ISAFP	YESAVI	NGIR	VLKTSD	NNCWVNA
Human corona 229E pol 1ab (1668 Murine hepatitis pol 1ab (1926) 文述CVE	YKEDLS	OYYC	BSGKY	MKPI	TRADER	TEEKVOG
Consensus (195) VKKVE	YAPDLS	OYYO	D SKYY	T T		TEEKVDG
							- Section 52
(1996) 1990	2	000	. 2	010		202
vian infectious bronchitis pol 1ab (1510							
bovine coronavirus pol 1ab (1920)) YYNYE	LUCHT	Chi	MAKIGE	DSSK	EEVEYR	VALUATA
Human corona 229E pol 1ab (170)	CIAL	YSKPHI	ISQC	LDAAWN	KEVL	GDEF	MAPVYYV
Murine hepatitis pol 1ab (1969) YTNET	LVCHS	ABKI	NAKUSI	DCNS	PHREXK	ITEWPTA
Consensus (1996							VTEWPTA
							 Section 53
	2029		2040		2050		206
vian infectious bronchitis pol 1ab (154) LTEK	RGIKST	MDFF	SKDGF	XKLT	PDTDEN	SKAPWYY
bovine coronavirus pol 1ab (1959) GDVVI	ATODE	CVKE	ERGOI	EGK E	VIWLSH	EOASLNS
bovine coronavirus pol 1ab (1959) Human corona 229E pol 1ab (1749)) RLMKO	DKGDAI	DEL	KLSKY	LANEA	OVOLEH	YSSCHEC
Murine hepatitis pol 1ab (2004)	I) GBWW1	ASDDE	IVER'	Sect.	PECKP	VVWLGH	CEASLKS
Consensus (202)) GDVVI	ASDDL	VSR	SKGCI	PFGKP	VVWLEH	E ASL S
							- Section 54
	3) 2068		2080		2090		210
vian infectious bronchitis pol 1ab (158	B) VLDA	SEKATI	IVEG	ANFEV	SHPNY	YS	
bovine coronavirus pol 1ab (199	B) TYPNI	PLLVD	ENKF	VIKEDI	DVDDG	G	
bovine coronavirus pol 1ab (199 Human corona 229E pol 1ab (178	5) AKEKI	SVASE	VIAL	CASTKI	RDGVQ	VG	
	at the second	In categoria	Market Mark	TEXT IN THE	z offilm m	DECRUP	3 3 11 1 11 mc
Murine hepatitis pol 1ab (204)	5) TXEN	ALDIA AC		A'A TEAL	ADELL	DAGEVE	AMADAIG

(2407	1 2407			Section 5
(2107 avian infectious bronchitis pol 1ab (1615	2101	2120	2130	21
bovine coronavirus pol 1ab (1615 Human corona 229E pol 1ab (1812	,			KSLI
Human corona 229E pol 1ab (1812	,		·j	ISESDAK
Murino honolitic and data (1012)		·	YCV
Murine hepatitis pol 1ab (2082) PGADASAG	AGIAKEQKAC <i>i</i>	SASVEDOVVTE	VROEPSV
Consensus (2107)		Î	I SVI
				Section 5
(2146	2146	2160	2170	21
2146) avian infectious bronchitis pol 1ab (1620) bovine coronavirus pol 1ab (2034)	PTEWENAE	NEWKMGDETGO	VTMCLWPARUT	21
bovine coronavirus pol 1ab (1020) Human corona 229E pol 1ab (1817)	KEINIDET	SCVKKPFKUPT	CULUMNAEHI	NKPNLER
Human corona 229E pol 1ab (2034 Murine hepatitis pol 1ab (2121) Consensus (2146)	TKYYSB	VRSVDCDARTS	PARTITION OF THE	ATIVES DS.
Murine hepatitis pol 1ab (2121)	ADVENTED	THE WAY TO STATE OF THE PARTY O	DATOTERCHOS	HLUSGVA
Consensus (2146)	DIRETET	がとれたい日本版が手を は対象が応じる本版が手を	SVIVNDPTSET	KYVKSES:
(21.10)	DIKEIKL	MGAWL KIEC	SVIVNDPTSES	KLVKSLS:
(0405)	0405 0400		·	Section 5
(2185)	2185 2190	2200	2210	22
avian infectious bronchitis pol 1ab (1659)	NIAKKARV	RSSVATTQCGE	DIGKAATFIAD	KVGGGVV
Human corona 2005 - 14 to (2073)	DAXDAMRA	CRYVETANA	I SMAVNVETTR	KFIKFGM
bovine coronavirus pol 1ab (2073) Human corona 229E pol 1ab (1854) Murine hepatitis pol 1ab (2160)	AFSGPVDK	HYTKYDTAKK	SMYDGDREVKH	DISLISVE
Consensus (2185)	DVYDMFLT	GCKYVV TANK	LS VNSPTIR	KVTKECVO
				Section 5
(2224)	2224 223	0 2240		22
avian infectious bronchitis pol 1ab (1698)	TITOSEKGE			
bovine coronavirus pol 1ab (2112)	WST PHONT	NERETEDVENS	Perdy Wellands - a see	TA CHEST TOTAL
Human corona 229E pol 1ab (1893)	VVMVGGVV	D DUNGUEDED	MANAGERICA	TURTRALL
bovine coronavirus pol 1ab (2112) Human corona 229E pol 1ab (1893) Murine hepatitis pol 1ab (2199) Consensus (2224)	TUTBARTO	THE ANTANE WAS	TROTHUERROKE	FDFG DF L
Consensus (2224)	TUTDIKI	THE WASTERNAM	IK AK KY VC	XCAV KWFI
(TATITUD	TKD K F. A	TK AK KY YC	
(2263)	2002 00	70		Section 5
(2203) Vian infectious bronchitic not tob (4746)	2263 22	270 228	0 2290	23
vian infectious bronchitis pol 1ab (1716)	KMSPQFLK	TEMFFOREELK	ASVAS	
Human corona 229E pol 1ab (1932)	NEVIELLM	LESMETECKEA	VITTGDVKIMAK	APORTGVV
widinie riepatitis por rab (2238)	YCDSWIKE	RIDNKVIYTEE	VASKETFKLCC	DAEKNALO
Murine hepatitis pol 1ab (2238) Consensus (2263)	LF WIKE	SLDNKVIYTTE	VASKLT KL	LAFKNALL
				- Section 6
(2302)	2202	2310 23	20 222	
(2302) Vian infectious bronchitis not 1ab (1741)	2302	TELEVISION AND THE	20 233	0 234
(2302) Wan infectious bronchitis pol 1ab (1741)	2302 VASYKTVL	CKVVLATLLIV	FVYTSNPVMF	O 234 FGIRVLDP
(2302) Wan infectious bronchitis pol 1ab (1741) bovine coronavirus pol 1ab (2190) Human corona 229E pol 1ab (1971)	2302 VASYKTVLO FKWSVVARO	CKVVLATILIV JACELATIFUL	EVYTSNPVMF ÆNETYANVIF	0 <u>23</u> 4 FGIRVLDP SDFYLPKI
(2302) Vian infectious bronchitis not 1ab (1741)	2302 VASYKTVLO FKWSVVARO	CKVVLATILIV JACELATIFUL	EVYTSNPVMF ÆNETYANVIF	0 <u>23</u> 4 FGIRVLDP SDFYLPKI

(2341)	2341	2350	2360	Section 61
avian infectious bronchitis pol 1ab (1780)	EECSI.CO	Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	2300	2379
bovine coronavirus poi 1ab (2229)	ET PRESENT	CHA ONTEON	Moster and the state of the sta	KDSFDVLRYCA
Human corona 229E pol 1ab (2010) Murine hepalitis pol 1ab (2316)	LTCVPF	AND TWO II	POTA TATE TARE	ODAGEKNOAGE
Murine hepatitis pol 1ab (2316)	PEDEFE	The a simple with	COBEWNGYA	KSNEVKDDYCD
Consensus (2341)	FLPTFUC	I W KST	CASTACDIAON	TDLGYRSSECN
	FREIFVG	I W KST	F L TICD Y I	KDLGFK YCN
(2200)	0000			Section 62
(2380)	2300	2390	2400	2418
avian infectious bronchitis pol 1ab (1804)	DDFICE	THOKOSEH	LYKHAYSTEQVY	KDAASGFIFNW
bovine coronavirus pol 1ab (2268)	CS-LAUQFE	LAGEDMED	NYKALDV QYEA	DREAFVOYTGV
Monte repailed put (2000)	TOTAL PARTY OF STREET	SESSION DOMESTS IN	MVDATENTIAL	Diff. with the manager of the
Consensus (2380)	GSIICKLO	LAGFDMLD	NAKHIDAAÕHAI	DRRLS DY V
				Section 63
(2419)	2419	2430	2440	2457
avian infectious bronchitis pol 1ab (1843)	NWEYTVEL	LLEVKPVA	GFNI	CYCVKYLVLN
bovine coronavirus pol 1ab (2307)	TRIVELL	VSYALYTA	MENCETER	LITTWIFELEM
Murine hepatitis pol 1ab (2394)	FXLVVELV	GYSLYTV	CEXPTEUTTOWN	Car difference of the comment
Consensus (2419)	IKTATETI	IGYALYTA	FYPLF LILIQ	ILTTWLPELFM
				Section 64
(2458)	2458	2470	2480	2496
avian infectious bronchitis pol 1ab (1874)	STVIQTGV	CFEDMEVO	PYFSHFNFMGAG	DV DEGENORE STATE
Dovine Coronavirus por 1ap (2346)	T.STTH WS.	PARTITION LAND	ST. D'A MATERIA	Frank Marian San San San San San San San San San S
Widine riepaulis por rab (2433)	LETHRESA	RIFIVEVAN	MER PORFTER DESCRIPT	FATTER WAY TO THURS
Consensus (2458)	LSTLHWSV	RLLVWFAN	MLPAHVILRFYI	I ALIKVI L
				Section 65
(2497)	2497	2510	2520	2535
avian infectious bronchitis pol 1ab (1913) bovine coronavirus pol 1ab (2385)	QVIHILYC	KDVTHEVE	KRVARSNEOEVS	WIGGBROTUH
bovine coronavirus pol 1ab (2385)	FRHVANGE	SKPOLLFE	KRNRSLHVÆCS	PRECMIEVAD
numan corona 229E poi 1ab (2146)	VRHVEFG	F MAD DO TA	THE A STATE OF DEALERS	This is a state of the state of
Mullie repails por rab (24/2)	CRHVMYGE	SEPERMENT	KRNIS STRAFFCOL	DAZIET C C T ITTERES
Consensus (2497)	RHVIYGO	SKPGCLFC	KRNRSLRVKVS	TVGGMTRYYD
				Section 66
(2536)	2536	255	2560	2574
avian infectious bronchitis pol 1ab (1952)	MYPHISEVN	BERDGNOV!	PHENDYCUON	THE RESERVE
bovine coronavirus pol 1ab (2424) Human corona 229E pol 1ab (2185)	MANGGTG	HESKHOWN	Thebeverent	TOTAL AND THE
Human corona 229E pol 1ab (2185)	NAUGUSK	FCKKHREE	Unensycycler	THORNEDO
Murine hepatitis pol 1ab (2511)	MANGETG	ECVEKNOWN	INCHSUK DOM	TIPE V SKEELS
Consensus (2536)	VMANGGTG	FCKKHOWN	CINCDSYKPGNT	TUDEUX ADEC
(2000)		- C.L.LIIQWIN	LINCOSIKPGNTI	TIPEVAADLS

2500 2500
avian infectious bronchitis pol 1ab (1991) E KET BEINK FIRA YEVAN FORACL VODE FVINKYKAART PGKT bovine coronavirus pol 1ab (2633) KOLIN REGO BOVS HIT NIT DYKKOVEC YMR RYSK MOGENT HUMAN CORONA 229E pol 1ab (2224) NITSTIN VOLK POS WYSK MIT DYKKOVEC YMR RYSK MOGENT HUMAN CORONA 229E pol 1ab (2224) NITSTIN VOLK POS WYSK MIT DYKKOVEC SMR. RYSK MOGENT HUMAN CORONA 229E pol 1ab (2650) KELKR PVQ PT D A YHT V TEVK V V V GC PMR LYSK BOGGR YMR RYSK MOGENT HOLD WAS AND SECTION AS S
Human corona 229E pol 1ab (2244) NITSTNIVONED PROVINCENCE YMR FYREDCERTS Murine hepatitis pol 1ab (2550) KETREMNIP DS FESTIVATOR MURINE HERGEYR ZSCETFWEYT Murine hepatitis pol 1ab (2550) KETREMNIP DS FESTIVATOR MURINE HERGEYR ZSCETFWEYT (2514) 2614 2620 2630 2630 2840 2660 avian infectious bronchitis pol 1ab (2030) SASSAVKCFSYTDTENKAWFLKEADKCEQLS PGFTVCN bovine coronavirus pol 1ab (2630) FDYNSTETTY SMALRS WKGY PMHVVWHAP
Murine hepatilis pol 1ab (2224) NITSTNIVOTE GPROVINCIONE PRINCIPER SCETTRICTS Consensus (2575) KELKRPVQPTD A YHTVTEVKQVGCSMBITYERIOGORY Consensus (2575) KELKRPVQPTD A YHTVTEVKQVGCSMBITYERIOGORY Section 68 2610 2630 2640 2650 avian infectious bronchitis pol 1ab (2030) SASSEVKCFS YTDFTEKRAWFLKEADICS OT SARGETIVCH bovine coronavirus pol 1ab (2202) DDVD) STFTEY NAME INKVKGVETHIVVVVING Human corona 229E pol 1ab (2263) FDTTEBKAGCKEVFKNCNYLEDFTYFTHOMICS
Consensus (2575) KELKRPVQFTD AYHTVTEVKQVGCFMRLFYERBGGRV3 Consensus (2575) KELKRPVQFTD AYHTVTEVKQVGCFMRLFYERBGGRV3 Section 68 (2514) 2614 2620 2630 2640 Avian Infectious bronchitis pol 1ab (2020) SASSEVKCFSVTDFTCKRAWFLKEADKCE QTSREGFTVCN Human corona 229E pol 1ab (2020) FDVVASEFTDYSMRLFISVKGVFMMVVVVVV Human corona 229E pol 1ab (2030) FDLTEFEXBCCKEVFVKN CONVLIDE FTUTINING
2614) 2614 2620 2630 2640 2650 2650 2650 2650 2650 2650 2650 265
Section 68 avian infectious bronchitis pol 1ab (2030) SASSENVKCFS WTD FTCK KAWFLKEAD KCEOLTS DEFIVED bovine coronavirus pol 1ab (2803) SASSENVKCFS WTD FTCK KAWFLKEAD KCEOLTS DEFIVED Human corona 229E pol 1ab (2263) FD TTEB KWSCKEV FKN CNYL DEFTWF NN G Murine hepatitis pol 1ab (2889) DDVNAST WTD MN GIDERS KYKCV BT THAVWYEN G Consensus (2614) DDVNASL FVDMS LHSKVKGV BT THAVWYEN G CONSENSUS (2614) DDVNASL FVDMS LHSKVKGV BD LHVVVVEN D Section 69 avian infectious bronchitis pol 1ab (2669) TQSAHALEEAKNALEV KYYCK WTD LHVVVEN D bovine coronavirus pol 1ab (2635)
avian Infectious bronchitis pol 1ab (2030) SASSEVIKCESTED PERKANFILKEALIKCE CITSIBGE FIVEN bovine coronavirus pol 1ab (2030) SASSEVIKCESTED PERKANFILKEALIKCE CITSIBGE FIVEN bovine coronavirus pol 1ab (2502) DOVINS EFFET PYSMELBSKYKGV POMHVYWY BENEVIKEN BOVING CORONAVIRUS POLITIC BOVING CITSIBGE FIVEN BOVING CORONAVIRUS POLITIC BOVING CORONAVIR
avian infectious bronchitis pol 1ab (2030) SASSINVICES OFFICE AMELIKEAD KCECTSABG FIVED bovine coronavirus pol 1ab (2502) DEVIN BETENDY NHLESKY KGV POMINVI WEST OF VALUE TO THE WAY OF THE
Human corona 229E pol lab (2263) FD_TESEMSC KEV FKN — CNYLDD FTYFVN FG — Murine hepatitis pol lab (2589) FD_TESEMSC KEV FKN — CNYLDD FTYFVN FG — Murine hepatitis pol lab (2589) FD_TESEMSC KEV FKN — CNYLDD FTYFVN FG — Section 69 Consensus (2614) DDVNASLFVDMS LLHSKVKGVPDLHVVVVEND — Section 69 (2653) 2653 2650 2670 2670 2680 2690 avian infectious bronchitis pol lab (2635) — FDARATER VALVE FEBELLY DANTED VVENDOWN FOR FROM THE CONTROL OF TRANSCORP PROVINCE FEBELLY DEN TARANTE Human corona 229E pol lab (2933) — TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FOR TRANSCORP PROVINCE FEBELLY DEN TARANTE FUNDOWN FUND
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Murine hepatitis pol 1ab (2689) DEVENTI DIAGNOCKEV KN CNVLID PTTYFNN G CONSERVATOR OF THE CONSERVATOR
Conseisus (2014) DDVNASLFVDMS LLHSKVKGVPDLHVVVVEND Section 69 Avian infectious bronchitis pol 1ab (2069) TQSAHALEEAKNSALVYKGYLCKETTILIDQAHYEQHVVE bovine coronavirus pol 1ab (2053) TOKANFILIN AVETA SEFRULIM OKNITTAN TO
Conseisus (2014) DDVNASLFVDMS LLHSKVKGVPDLHVVVVEND Section 69 Avian infectious bronchitis pol 1ab (2069) TQSAHALEEAKNSALVYKGYLCKETTILIDQAHYEQHVVE bovine coronavirus pol 1ab (2053) TOKANFILIN AVETA SEFRULIM OKNITTAN TO
2653 2650 2670 2680 2690 avian infectious bronchitis pol 1ab (2659) TQSAHALEZAKNALTAYAÇYICKETATI TQAHYEQTIYVE bovine coronavirus pol 1ab (2635) TAYARIYAN AYEYAÇSIFRALIMYEKINI TÜAN TE Human corona 229E pol 1ab (2293) TAYARIYAN TAYARIYA
avian Infectious bronchitis pol 1ab (2069) TQSAHALEEAKNISLINIKGY CRETATIOOATINO DIVISE bovine coronavirus pol 1ab (2935) TOKANTIN AVETA SEFRULIMOKNITTANTE Human corona 229E pol 1ab (2933) TOKANTIN AVETA SEFRULIMOKNITTANTE
avian infectious bronchitis pol fab (2069) TQSAHALEEA KINSLEYYA GYECKETLIJI QAHYEQEVVE bovine coronavirus pol fab (2535) — ——ADVANTEN AVETTA SEFENDLIMVAKNI TTANTO Human corona 229E pol fab (2933) ——TNIVOTAVESTA
Human corona 229E pol 1ab (2293)TRIVERVENT TO SEE FROM THE AVERAGE FROM THE TANTE
Consensus (2653) ADKANFLNAAVFYAQSLCRPILMVDK LITTLNVG
Section 70
(2692) 2692 2700 2710 2720 2720
avian infectious bronchitis pol 1ab (2108) PVS - SSWITAKYCS TOSERY I SWEET TO BEEN I SWEET TO BEEN I SWEET TO BEEN I SWEET TO BEEN I SWEET TO BEEN I SWEET TO BE T
DOVING CORDINAVIOUS DOI 12D (2570) THE THE PROPERTY OF THE PRO
Widnie nepalits por lab (2007) LSVSRFMEDTY AND STREET, BY DESCRIPTION OF STREET
Consensus (2692) SVSKTMFDLYVDTLLSIFDVDKKSLNA I AH.SIK
Section 71
(2731) 2731 2740 2750
avian infectious bronchitis pol 1ab (2134)VKACTIED AND THE PROPERTY AND AND AND AND AND AND AND AND AND AND
bovine coronavirus pol 1ab (2609) STOECKVDDTFLSCARKSCSIDSDVDTKCLATSVMSAVS
Human corona 229E pol 1ab (2352)MSLAECERRAIGLSISDHEPISATSNOHR
Murine hepatitis pol 1ab (2696) GVOLE QVINDTELIC CARRACCATOS SYNCIAS TIKS VINSAVN
Consensus (2731) G QI VLDTFISCARKKCAIDSDVDTKFITDSVMCAVN
(0770) 0770 Section 72
(2770) 2770 2780 2790 2808
avian infectious bronchitis poi 1ab (2162) HDWDYTGDGFTNVTPS GIDTGKITPRSRCFLINADAS I
Human occomination pol 1ab (2044) RG DEL HUD BC CONST. VID THE RG - DNI LAAS LGVITTONS KKE
bovine coronavirus pol 1ab (2848) AGUELI OBSCNAL VESTICI DTOKLT RADRICINIANAN SI Human corona 229E pol 1ab (2880) CDVILL SID SEN PONSAN ARP BERLISAY CITACONRAGISM Murine herafilis
Consensus (2770) AGVDLTDESFNNLVPSYLK DKIVAADLGVLIQN AKH

							Section 7	3
. (2809) 2809		2820		830		28	
avian infectious bronchitis pol 1ab (2201) ANLRYK	NAP	PVVWKE	SELIE	TEDSC	LKYLI	SATVK	3.6
bovine coronavirus poi 1ab (2686) VOGNYA	KIAGV	SCIWSV	DAFNO	L'SSDF	онкт к	KACCKI	ρź
Human corona 229E pol 1ab (2419) VNANEL	TKDOT	PIVEHA	KDFNS	LSAEG	RKYIV	KTSKA	ΚĒ
Murine hepatitis pol 1ab (2773) VOANYA	KAANV	ACIVEV	BAEN	EADL	OHRLR	KACSK	ijŹ
Consensus (2809) VNANVA	KAANV	PCIWSV	DAFNO	LSAD	ÖKYLR	KAC K	řc
					-		Section 7	14
(2848	3) 2848		2860		2870		28	38
avian infectious bronchitis pol 1ab (2238) VREFI	KSGAK	OVIACE	TOKLI	VEKKA	GGLVS	GTFKC	ĒΈ
bovine coronavirus pol 1ab (2725) LKLKLT	ANKO-	-MANVS	VLTT	FRIEG	AVES		
Human corona 229E pol 1ab (2458	B) TELL	INENO	AMTOTE	ATSIV	AKOGA	GDAGH		
Murine hepatitis pol 1ab (2812) EXIKET	YNKO-	EANVE	PLTT	PELKE	SAVES]	
Consensus (2848) LKFKLT	YNKQ			FSLKA			
·							Section 7	75
(2887	7) 2887		2900		2910		29)2
avian Infectious bronchitis pol 1ab (2277 bovine coronavirus pol 1ab (2755) SYFKWL	LIFYI	HTACC	SGYY	MEVSK	SEVHP	MYDVN	S
bovine coronavirus pol 1ab (2755	i)	YEMXV	CELLSI	V.CHT	LWCLM	PIYTY	HKSDF	ō)
Human corona 229E pol 1ab (2490))	SLTWI	WLIJCGI	UCLI	PENTOF	FMPXF	MYDIV	S
Murine hepatitis pol 1ab (2842	2)	RMLQW	PVANÍ	TOFI	TWALM	PTYAV	HKSDM	٥١
Consensus (2887	r)						MYSDMS	
							Section 7	76
(2926	3) 2926		2940		2950		29	
avian infectious bronchitis pol 1ab (2316) LHVEGT	EVEDK	GVTRE	VPED	POPSHK	EVNEE	AEWGR	P
bovine coronavirus poi 1ab (2788	B) PVYASY	RVLDN	VIRD	SVED	FATK	PROFI	OWYES	ń
Human corona 229E pol 1ab (2523) FEGYDE	EYIEN	GOLKNI	EAPL	VRHV	EENFE	DWHYAI	ĸ
bovine coronavirus pol 1ab (2788 Human corona 229E pol 1ab (2523 Murine hepatitis pol 1ab (2878	5) BIYASE	"VIDN	VLRD	SVTD	CFAVE	PNOFE	OWYES	m
Consensus (2926	PLYASE	KVIDN	GVLRD	SVED	CFANK	FENFO	OWYES	ΤÏ
							Section 7	
(2965	5) 2965 2	970	29		299		30	
avian infectious bronchitis pol 1ab (2355) DNSRNC	PIVTA	VIDGDO	TVAT	VPGFV	SWVMI	GVMFI	H
bovine coronavirus pol 1ab (282)								
Human corona 229E pol 1ab (2562	2) GFTPLN	K-Q		PIVV	VSEIV	NIVAC	LPSNM	Y]
Murine hepatitis pol 1ab (2914	1) GEAYYP	NSKA-	CP	PVVAV	I DQDT.G	HTLEN	VPTTV	G.
Consensus (2968	5) GLSYY	NSMA	CP	VAV	GVÕDIV		VPT V	
							- Section 7	78
		,3010		3020		030	30	
avian infectious bronchitis pol 1ab (2394	4) TOTERS	PWYIP	TWFNRI	IVGY	FODSII	TEGSE	YTSIA	Ť
bovine coronavirus pol 1ab (286)	2) YGYHVI	HEITH	ALSAD	VOCY	PHSOI	SYSNE	YASGC	V.
Human corona 229E pol 1ab (259	4) VGKTLI	FTLQA	AFGNAC	SVCYD	IFGVTT	PEK	C	I
Murine hepatitis pol 1ab (294)	9) YGFHVI	HEITH	AFATO	VOCY	PHMOI	PYDNE	YASGC	v
Consensus (3004	4) YGFHVI	HFITH	AFANDO	SVOCY	TPHSOI	PY NE	YASGC	v

					- Section 79
(3043)	3043	,3050	3060	3070	308
avian infectious bronchitis pol 1ab (2433)	SARLL	YETASHTPO	LYCFNGDND	APGALPEG	STIPHRY
bovine corona 229E pol 1ab (2931) Human corona 229E pol 1ab (2627) Murine benatilis pol 1ab (2088)	SSACT	MEAMADGS	PORYGYTEGI	MONASLYS	SLVPHVR
Human corona 229E pol 1ab (2627)	TSAST	REEGLGGN-	NVYGYNTAL	MEGSLPYS	STOANAY
marine ricpatitis por rab (2500)	2070	HUMBLET	PHPYCMTICGV	MHNASITYS	CT A CHITTO
Consensus (3043)	SSACT	MLAAADGS	PNPYCYTDGL	M NASPYS	SIIPHVR
					- Section 80
(3082)	3082	3090	.3100	3110	312
avian infectious bronchitis pol 1ab (2472)	FOPNG	WRLIVPO	OILHTPY	PRUS DOVE	Hook Way
bovine coronavirus por tap (2940)	MINAMA	KGFTREER	TTP DOCT TO DO	DUDGALCUC	CAR CONTRACTOR
numan corona 229E por 1ab (2665)	KYDNG	METRI PRU	CHOCKCREEN	D III T Inim ve Sulos	10 77 7 m 12 77 78
Mulline nepaula por rab (3027)	N.L. A. 5 5	MCXTHEFE	MISEOTARUS	RITERIS NETWORKS	Water recent
Consensus (3082)	NLANG	NGVIRFPEY	L EGIVRVV	RTRSMSYC	RVGLCEE
					- Section 81
(3121)	3121	3130	3140		315
avian infectious bronchitis pol 1ab (2509) bovine coronavirus pol 1ab (2979) Human corona 229E pol 1ab (2004) Murine bozpitis pol 1ab (2004)	RPY	VSLNPQW	FNDEYTSKE	CVEDGSTV	REIMESM
bovine coronavirus pol 1ab (2979)	DEST	FNENGSW	NOUTYRELP	GTEGERDV	FDUITYCL
Human corona 229E pol 1ab (2704)	NAGVO	EGEDKWE 1	IDGRVAN	GYVOGTGL	MNUVENI
munite nepauls pur lab (3000)	#1 b 15 5 12	FUND N. BONDON	CHARD MACOUNT MAD	THE TRUTH THE TOTAL A	TAIR TOTAL CO. A.Y.
Consensus (3121)	DEGIC	FNFNKSWV1	UNDALESTE	GTFCGR V	FDLIFQI
					- Section 82
(3160)	3160	,3170	,3180		319
avian infectious bronchitis pol 1ab (2548)	STFF	GYNPN-EYN	IQLATMFLIL	VVVVLIFIA	MVENFOG
bovine coronavirus pol 1ab (3018) Human corona 229E pol 1ab (2740) Murine benatitis pol 1ab (315)	KOLAO	PVDELALT?	SSIAGALLA	AINATALA	YLIFIKR
Human corona 229E pol 1ab (2740)	SMFSS	SFSVAAMS	QILLNCALG	AFATFCCE	LVTEFRE
marine nepatitis per rap (3 ras)	COTAK	THE PARTY A	SSVAGALLA	TTVATEARY	YELD TAKE
Consensus (3160)	SGLAS	PVDF ALT	SSIAGAILA	VIVVLIFY	YLIKLKR.
				·	— Section 83
(3199)	3199	3210	322	:0	323
avian infectious bronchitis pol 1ab (2586)	KAYA	TTVETTML	WVINAFILC	MISYNSVI	AVILLYL
povine coronavirus pol 1ab (3057)	CDYT	STYFYNYIT	wevermmle	VEGVYPPL	SCVYAIC
Human corona 229E pol 1ab (2779)	GDLS	VGLCTVXV	VLLENVSYI	TONLVIM	IAYAILY
bovine coronavirus pol 1ab (3057) Human corona 229E pol 1ab (2779) Murine hepatitis pol 1ab (3144)	EGDYT	SALATIVATION	WOINERMIE	V FOVYPIL	SCEYACE
Consensus (3199)	FGDYT	SIVFINVI	WCINFLMLF	VFQVYPTL	SCIYAIF:
					Section 84
(3238)	3238	32	50 .32	260	327
(3238) avian infectious bronchitis pol 1ab (2625)	CMAST	VTSRNTV	MUCHTARRE	Cit it Tar mya'ri	A CONTICE
(3238) avlan infectious bronchitis pol 1ab (2625) bovine coronavirus pol 1ab (3096)	CYASL	VTSRNTV	MHCWLWFTE	GLIVETHE	ACCLLGE
(3238) avian infectious bronchitis pol 1ab (2625) bovine coronavirus pol 1ab (3096) Human corona 229E pol 1ab (2818)	CYASL FYATL FATES	VTSRNTVŽÍ YFPSEISVI LRYA-WI	MHCWLMFTE MHLOWLVMY WCARY TAY	GLIVPTWI GTIMELUIF	ACCLLGF CLLYISV
(3238) avian infectious bronchitis pol 1ab (2625)	CYASI FYATI FATRS FYTTL	VTSRNTV <mark>I</mark> YFYSEISV LRYA-WI YFPSEISV	MHCWLMFTF MHEOWLVMY WCAAYLLAY MHLOWLVMY	GLTVPTML GTIMPLUF ISFALWUL	CLL (ISV) CAW(ZFLA)

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(32	277)	3277		3290		3300	3316
avian infectious bronchitis pol 1ab (26	664)	HYMYTPL	FLWCYC	TTKN	TRKLYD	GNEFVENY	DLANKS .
bovine coronavirus pot 1ab (31	1351	VB	-NHAFL	VIDAV	DOT C'T	CHECK	THE PARTY AND ADDRESS.
Human corona 229E pol 1ab (28	854}	LTGL	-LPSEI	KLKV	STNEFE	DKFV	RCD AND COM
Munne nepatitis poi 1ab (32	222)	VS	-NHALV	LESY	REIST	EVESDOTE	EMALET
Consensus (32	277)	VS	NHALV	LFKY	CRKLGT	GVRSVGTF	EEMALTTE
							- Section 86
(33	316)	3316		.3330		3340	335
avian infectious bronchitis poi 1ab (27 bovine coronavirus poi 1ab (31	703) ·	VERGSEE	VANTER	G-D	CEE AVE	GAN ADTA	VECTCSEC
bovine coronavirus pol 1ab (34	168)	MITEDSY	CATIONICS	T.S.DW	ALTINE	CT MKVD	
Human corona 229E pol 1ab (28	8891	V. DMRSY	ERMAN	TSPE	CT.KS D	ASKADARA	VENT D NE
Murine hepatitis pol 1ab (32	255)	MERKESY	CONTRACTOR	VED V	A POT DE LA	STANKER	
Consensus (33	3161	MITKDSY	CKLKNS	TSDVI	KENRYT.	SIVMENT	VECKNDA
	,		OKZKK	71000	KI WKI D		- Section 87
(35	355)	3355 336	šo.	3370	0	3380	339
avian infectious bronchitis pol 1ab (27	741)	WAT OWED	Kings vs	55 C C C C	No of	23300	339
bovine coronavirus pol 1ab (3)	207)	AMPANA	Sove to	MOTOR	DAMAGE.		I SIGVER
bovine coronavirus poi 1ab (32 Human corona 229E poi 1ab (29	9281	DVBCLCV	AVITA	MT.D.	SID DUN _		TO VOLUME
Murine hepatitis pol 1ab (32	294)	AVERDAC	SOLUTION.	METER	HANGEN		V G I I G T G T I
Consensus (33	355)	DYREACC	AOI. AKZ	MDTR	S NNC	DIT VEDDE	2011年11日の日本日本日本日本日本日本日本日本日本日本日本日本日本日本日本日本日本日本
		211121100	TIQUITICE.	IIIDIE.	J 1010G	DIBITEEL	- Section 88
(33	394)	3394 3	400	34	10	3420	343
avian infectious bronchilis pol 1ab (27	779)	CSGFKAL	VSHSST	MIKE	EVSVS	RGNNEWOT	CTOTELY
bovine coronavirus pol 1ab (32 Human corona 229E pol 1ab (29	246)	USGIVEM	VNETSE	W. Per	CVSVT	NMT TO	HIDHKYN
Human corona 229E pol 1ab (29	965)	DAGLRICH	AOPSGE	Z.K.	VERVE	GNTURNGL	ERICH TATE
Murine hepatitis pol 1ab (33	333)	OSCIVIM	VSPTS	W. P.2	LICATE	NMTSACO	DEKV
Consensus (33	394)	QSGIVKM	VSPSSE	VEPC:	IVSVTY	GNMTLNGL	LGDKVYC
							- Section 89
(34	433)	3433	3440		3450	3460	347
avian infectious bronchitis pol 1ab (28	818)	FRINIGK	FSGDQV	INDVL	HANNH	EFEVTTOH	G——议工道的
bovine coronavirus pol 1ab (32	2851	DP HICH IN Q	A-C DMON	THE WATE	TTTCDT	THE STATE OF THE S	POSTERNI
Human corona 229E pol 1ab (30	0041	PRHATAR	M - TTSIZ	TWYD	HEVETM	DIUNDOWN	COM A PROCE
Murine nepautis por Tab (33	31Z)	PREPLUS	SADMUL	PDYP	NLUCRV	TSSDFCMM:	SGRMSLIT
Consensus (34	433)	PRHVICS	ASDMT	PDY I	NLLCRV	TSSDFTVI	SGRLSLTY
							- Section 90
10.		3472	3480		3490	3500	351
	oees	VESRRICK	AVETLO	TAVA	TAEUPK	YKEIKANC	TO DESTRIBA
avian infectious bronchitis not 1ab (28	ردده						Ethick Coll and the
avian infectious bronchitis pol 1ab (28 bovine coronavirus pol 1ab (33	324)	MSYOMOG	CMLVL	VTLO	SRIDK	YTEGVVKP	SOTETVL
avian infectious bronchitis pol 1ab (28 bovine coronavirus pol 1ab (33	324)	MSYOMOG	CMLVL	VTLO	SRITK MHT R	YTEGVVKP HSPRTEKS	GDTETVL SEGFNIL
avian infectious bronchitis not 1ab (28	324) 042)	MSYOMOS VGATMHG	OMLVE'	(VSQT	MHTER	HSTRTLKS	SEGFNIL

				Sectio	n 91
(3511)	3511	3520	3530		3549
avian infectious bronchitis pol 1ab (2894) bovine coronavirus pol 1ab (3363)	ALGETVVI	LYPUTMRS	NGTIRASMI	AGASOSVOFNI	EKC
povine coronavirus pol 1ab (3363)	ATNEKPOO	ATHV.TMR	SYNIKGSF1	CCSCCSVGYVL	MGC
Murine nepatitis poi 1ab (3450)	ANNERPO	TA FIRM TOTAL	CHATRCERA	Contract Contract	more
Consensus (3511)	AYNGKPQO	SAFHVTMRS	SWTIKGSFI	CGACGSVGYVL	ĞG
				Sectio	n 92
(3550)	3550	3560	3570		3588
avian infectious bronchitis pol 1ab (2933)	VORFE	HIRLPNAI	HTGTDLMER	EVGGEVTERNA	Sec. 10.00
bovine coronavirus pol 1ab (3402)	CKLYM	QUEL'STGC	HTGIDENSE	FERRYKIANUV	10 TO 10
bovine coronavirus pol 1ab (3402) Human corona 229E pol 1ab (3120)	ENERVANI	OIEGGS	HVISSTROCK	MEDGRENODNE	200
Murine hepatitis pol 1ab (3489)	STREET	OUL STG	HT-ABBSC	FIVEDVENANTA	7 6
Consensus (3550)	VKFVYME	OLELSTGO	HTGTDF GD	FYGPYKDAQVV	外が形の
				Section Section	v os Sme
(3589)	3589	3600	3610		-
avian infectious bronchitis pol 1ab (2972) bovine coronavirus pol 1ab (3441)	D DISNITUEN	1 THE R. LEWIS CO., LANS.	THEY PE	DOT DESIGNATION OF	3021
bovine coronavirus pol 1ab (3441)	VOOVTRE		WALLEY.	L 2 P L M THE SLLI	VIDV
Human corona 229E pol 1ab (3159)	SANOMICT		SOUTH STORY	UNWEVOSUK	ÇŞν
Murine hepatitis pol 1ab (3528)	VADETOR	A STATE OF THE STATE OF	TT CMD	CIWMIKGER	LEV
Consensus (3589)	VODVIOT	PHYPARAMETER VA	DIT N	CNWFLOSDS	Car
	.521151	MVVAWIIA	WIDN	Section	
(3628)	3628	3640	3650		
avian infectious bronchitis not tab (3011)	TO VOLKERY C	DO-TO	0000 m n m restric	MEET THE TOTAL OF EVER	3666
avian infectious bronchitis pol 1ab (3011) bovine coronavirus pol 1ab (3474) Human corona 229E pol 1ab (3192)	P C TO THE SAME	E TOTAL	ODTAGIA	ST CRADACKTR	RT
Human corona 229F not tab (3192)	EUN TEN	A THE COUNTY	CODDITION	an craptility	AAT
Murine hepatitis pol 1ab (3561)	PERMIT		DEDATE	AKE CYLKEL	HAL
Consensus (3828)	PDEMINAT	PAROE DOLL	ADDAMA	AMTGVSVEKLL	AA J
Generalia (802a)	EDENVMAL	DONGESAIK	ADLVIDALA		
(3667)	2007			Section	
avian infectious bronchitic not tab (2050)	3007	3680	369	0	3705
avian infectious bronchitis pol 1ab (3050)	MAKMSOMC	O DP THEOY	NEEDELT DE	STENUTGEVE	Q&-
bovine coronavirus pol 1ab (3513) Human corona 229E pol 1ab (3231)	WKTK MERC	HROTM. SC	SFEDELTES	DAAOFTVCPK	Q K
Musino honotitio nel 1et (3231)	CATIONER	EKOTPGAS	SLNDEFSIN	E VK MFGVNL	QS-
Murine hepatitis pol 1ab (3600)	REFEEE	PEROTIES	VERGETTES	DAXOCTACAKE	JS R
Consensus (3007)	KVLNSGFQ	GEKÖILGEC	SLEDELTPS	DVYQQLAGVKL	
				Section	n 96
(3706)	3706	372	20 ,3:	730	3744
avian Infectious bronchitis pol 1ab (3088)	-SFVRKAT	S-WEWSRC	VLACELEVE	CAIVLFTAVPL	KFY
povine coronavirus pol 1ab (3552)	RTRLVKGI	VERMARA	FIRSCATTON	DIVERTMAKEMAYYM	TI AT NO
wurine nepatitis poi 1ab (3639)	RTRVIKGT	CCWILAST	FLEGSTISA	EVKUTMEMYVT	THE
Consensus (3706)	RTRVIKGT	CWILAST	FLECETIST	FVKWTMFMYVT	TI EN

					- Section 97
(3745)	3745 3750	37€		3770	3783
avian infectious bronchitis pol 1ab (3125)	VYAAVILIA	MANLEISFT	VENVMAY	DTELLET	TITVIIG
bovine coronavirus pol 1ab (3591)	ISTITECAL	VISLAMLL	V. HKHLYI	TMYITEA	LIFTLLYN
Human corona 229E pol 1ab (3307)	TPEMILLY	ESLCLTEV	VKLKVLFI	OVELIE	ELVARIO
Murine hepatitis pol 1ab (3678)	DGVTLCAL	FVSFAMLL	ILHKHLYI	TMYINE	LCTLIFYT
Consensus (3745)	L ITICLLO	LVSFAMLL	VKHKHLYI	TMFILPY	LITLIYN
					- Section 98
(3784)	3784 3790	3 (3	800	3810	3822
avian infectious bronchitis pol 1ab (3164)	VCAEMPFI	NTLISOVV	TFLSQWY	PVVEDTA	AV PWMELP
bovine coronavirus pol 1ab (3630)	NYLVVYKO	FRGYVYAW	LSYYVPS	EYDYTDE	VIYCMLL
Human corona 229E pol 1ab (3346)	NCAWDEHV	CKVEAEKFD	YNVSVMO	MIOGEV	FFT CIEFM
Murine hepatitis pol 1ab (3717)	NATANAXXO	FRGUAYAW	ESHFUPA	DYLYMDE	VIXCVVI
Consensus (3784)	NATAAAKÖ	FRLIAYAW	LSVSVPA	DYTY DE	VIYGLLL
					- Section 99
(3823)	3823 38	30	3840	,3850	3861
avian infectious bronchitis pol 1ab (3203)	EVEYTAFKO	суоссумия	FNTSLLM	XOFVKLO	FVIYTSS
bovine coronavirus pol 1ab (3669)	LIGMVFVT1	RSINHD	<u>-</u>	FSFIME	GRVISVV
Human corona 229E pol 1ab (3385)					TYNESTE
Murine hepatitis poi 1ab (3756)	TWAMVEVED	ARS INHO			GRLVSLV
Consensus (3823)	LVLMVFVT	LRSINHD]		GRLVSLV
					Section 100
(3862) avian infectious bronchitis pol 1ab (3242)	3862	8870	3880	3890	3900
avian infectious bronchitis pol 1ab (3242)	NTLTAYEE	SNWEHFFEL	VHTTVLA	IVSENSL	GLFVFKC
povine coronavirus por rab (3699)	SHWYMGSNI	DE PETT PW	APTER	LWITTAES!	MAAAKVIA
Human corona 229E pol 1ab (3409)	AVLYTALY	SYDYVSLLV	MLLCAISI	EWYIGA	IFRICRE
Murine hepatitis pol 1ab (3786)	SMWYFGAN	PEERVINFI	TOTEGRA	CWITMES	ATAKVIA
Consensus (3862)	SLWY GSN	TEEEAPPPF	MSLFGTY		
					Section 101
(3901)	3901	3910	3920	-	3939
avian infectious bronchitis pol 1ab (3281)	AKWMLYYCI	IATYLNNYV	LMAVMVN	ILGWICT(PEGDYWW
bovine coronavirus pol 1ab (3738) Human corona 229E pol 1ab (3448)	KMAYANAT.	TTIDI POIK	IATACAT	PEGETI SC	EWGLESIL
Human corona 229E poi 1ab (3448)	GVAFEPVE	VSYFDGVK	LATITLAN	LTHENSEN	4 X G LU Y W
Murine hepatitis pol 1ab (3825)					
Consensus (3901)	KMTATUAT	YFTYIPQIK	TALL AL		
					Section 102
(3940)		,3950	3960	B.1 198	3978
avian infectious bronchitis pol 1ab (3320)					
bovine coronavirus pol 1ab (3777)	MNSLIFRMP.	USV NYKIS	VOELRYM	VANGLEP	KNSFEAL
Human corona 229E pol 1ab (3487)	LERFCKCT	CAN DECA	PAEFKYM	VANGLNA	RNGPFDAL
Murine hepatitis pol 1ab (3864)	L. PS LERM P	LUYYNYKI	VOEDRAM	NANGLRPI	PRNSFEAL
Consensus (3940)	INSIFRMT	LGVYNFKIS	VQELRYM	NANGLRPI	PKNSFEAL

					Secti	on 103
(3979)	3979	,3990	male terrories	4000	****	401
avian infectious bronchitis pol 1ab (3359)) STNILI	QGIGGDRI	LPIATV	CARLSTV	CTTLLL	OLL
bovine coronavirus pol 1ab (3816	MINEKI	LGTGS VPI	IEVSQF	CSKLOSVK	CANAVIT	NOL
numan corona 229E poi 1ab (3526)) FISEKLI	MATHEMATIN	TRUSTY	TO LOS TO STATE	CORP STREET FOR	
wunne nepatitis poi 1ab (3903)	MINERE	DGDGGVPV	LEVSOI	OSBUTTOUR	CANTENT	NOTE:
Consensus (3979)	MLNFKL:	LGIGGVRV	IEVSTV	QSKLTDVK	CTNVVLI	NCL
					Section	on 104
	4018	40:	30	4040		405
avian infectious bronchitis pol 1ab (3398)	KINVEA	ESEMHVYI	VETSTR	TASDING	ECMBARS	7 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
bovine coronavirus pol 1ab (3855)	HLHVAS	SECONO	STITUTE	PAPSITE	VAFEKTIZ	6
bovine coronavirus pol 1ab (3855) Human corona 229E pol 1ab (3565)	NMNIAS	EVALE	VEMENK	NICOPE	TAORT	n is
widine nepadas por lab (3942)	HUHLAS	HASTE IN OUR C	STATE	THE DIE CHAPTE	AZ TO TOTAL TO T	CO. Tarte
Consensus (4018	HLNIASI	NSKLWOYC	VTLHNK	TLATSDIG	VAFDELL	OLI
					Section Section	on 105
(4057)	4057	4	070	4080		
avian infectious bronchitis pol 1ab (3437)	TEECTD	START	- Higher	OND THE DE	militaria.	CHAIL CO
DOVIDE CORODAVIRUS DOI 14h (3894)	13 TO 13 TO 15 TO 15	ALT TETT COLCUT	IDICY TOTAL TO	CHEST PROPERTY.	The state of the s	A 40 4 4 4 4 4
Murine hepatitis pol 1ab (3981)	MANUTANT	AWKEZH	ACTECI	CONTRACTOR AND	OT LUDY!	の心臓
Consensus (4057)	VIFAND	AAVDSKCT	STERV	CDDATADA	THE	京都県
			OIBEV	CDDIBRON		on 106
(4096)	4096		4110	4120		440
milan infantiaria hannalitta ant det da emor						
avian iniectious pronchitis poi 1ab (3470)	HTPREA	THREE IN	OKE WILLIAM	Dennecum	ANDER A	T WELL
avian infectious bronchitis pol 1ab (3470) bovine coronavirus pol 1ab (3933)	AT ME TO SEP CUTTO	CONTRACTOR OF STREET	T TOTAL TOTAL	DSKNGGVT	OCELAAY	Riva
Human corona 229E pol 1ab (3637)	NMASEV.	IN FUNKKN	LDEARS	DSKNGGVT SCSAN	ODOLKOT	REZ
Human corona 229E pol 1ab (3637)	NMASEV.	IN FUNKKN	LDEARS	DSKNGGVT SCSAN	O OLKOI PHILKOI	BKA KEAI
Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020)	MASEV GMP4EV NMASEV	EN EVAKEN ATETAROE EVSTAKKN	LDERRS YENAVA LDRAKA	DSKNGGVT SSAN NGSS SGSAN	O OLKOI POIIKOI Odolkoi	BKA KFM BKA
Human corona 229E pol 1ab (3637)	MASEV GMP4EV NMASEV	EN EVAKEN ATETAROE EVSTAKKN	LDERRS YENAVA LDRAKA	DSKNGGVT SSAN NGSS SGSAN	OOOIKOI PRIIKOI OOOIKOI	BKA KEAI EKAI
Dovine coronavirus por 1ah (3933) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096)	MASEV MASEV NMASEV	ATHTARQE EYELAKKN	LDEERS YENAYA LDEARA YDEARA	DSKNGGVT SESAN NGSSI SGSAN SGSAN	O ØLKO PDILKOI O O LKOI QQQI KQI ——— Sectio	EKA EKA On 107
Dovine coronavirus por 1ah (3933) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096)	MASEV MASEV NMASEV	ATHTARQE EYELAKKN	LDEERS YENAYA LDEARA YDEARA	DSKNGGVT SESAN NGSSI SGSAN SGSAN	O ØLKO PDILKOI O O LKOI QQQI KQI ——— Sectio	EKA EKA On 107
Dovine coronavirus por 1ah (3933) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096)	MASEV MASEV NMASEV	ATHTARQE EYELAKKN	LDEERS YENAYA LDEARA YDEARA	DSKNGGVT SESAN NGSSI SGSAN SGSAN	O ØLKO PDILKOI O O LKOI QQQI KQI ——— Sectio	EKA EKA On 107
Dovine coronavirus por 1ah (3933) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096)	MASEV MASEV NMASEV	ATHTARQE EYELAKKN	LDEERS YENAYA LDEARA YDEARA	DSKNGGVT SESAN NGSSI SGSAN SGSAN	O ØLKO PDILKOI O O LKOI QQQI KQI ——— Sectio	EKAC
Dovine coronavirus por lab (3932) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096) (4135) avian Infectious bronchitis pol 1ab (3509) bovine coronavirus pol 1ab (3969) Human corona 229E pol 1ab (3672)	GEFLEN GEFLEN NAARLY NMPSFVI 4135 41 TAKSVI TAKSA	HOUNEN ATHTAROE EYELAKKN 40 FDEDLATO (EEDRAYA	LDEERS ZENAVA LDEARA YDEARA 4150 KE EDS	DSKNGGVT SESAN SGSAN SGSAN 4160 ER METN	Q Q K C I P I I K Q I P I I K Q I K Q I K Q I K Q I C Q I K Q I C Q I K Q I C Q I K Q I C Q I K Q I C Q I K Q I C Q I K Q I C Q I K	EKA EKA On 107 417 DRB
bovine coronavirus poi 1eh (3932) Human corona 229E poi 1eh (3637) Murine hepatitis poi 1eh (4020) Consensus (4096) avian infectious bronchitis poi 1eh (3509) bovine coronavirus poi 1ab (3889) Human corona 229E poi 1eh (3672) Murine hepatitis poi 1eh (4056)	MASSIVITY OF THE PROPERTY OF T	INGUAREN ATHTAROE EYELAKKN 40 DECLATO GEEDRATA FORESSA	LDEERS YENAVA LIDEARA YDEARA 4150 KELDSIN RELESIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN	DSKNGGVT	O O KON PDILKOT O O EKOI QOQIKQI Sectlo IZMAIVI UEARIN	EKA EKA On 107 417 DRB DKK NRK
Dovine coronavirus por lab (3932) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096) (4135) avian Infectious bronchitis pol 1ab (3509) bovine coronavirus pol 1ab (3969) Human corona 229E pol 1ab (3672)	MASSIVITY OF THE PROPERTY OF T	INGUAREN ATHTAROE EYELAKKN 40 DECLATO GEEDRATA FORESSA	LDEERS YENAVA LIDEARA YDEARA 4150 KELDSIN RELESIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN KELDSIN	DSKNGGVT	OBOTKOT POTENCIA OTOTICA OTOTICA Section THE ASSOCIATION THE A	EKA EKA On 107 417 DRR NRK DKK DKK
bovine coronavirus por 1ab (3933) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096) evita infectious bronchitis pol 1ab (3509) bovine coronavirus pol 1ab (3692) Human corona 229E pol 1ab (3672) Murine hepatitis pol 1ab (4056) Consensus (4135)	MARCH GEPTEN NMARCH NMPSFVI 4135 41 FTAKST FTAKST NIAKST NIAKST	ALUTEROE EYELAKN 40 EDROLATO EDROLATO EERORAYA FORDRAYA	LDERKS MENAVA LDEAKA 4150 KELUS RELUS RELUS KELU	DSKNGGVT SCANG	OBOLKOL POLLEKOL OGOLKOL OGOLKOL Section LELARUT LELARUT LELARUT LELARUT LELARUT YKEARIN YKEARIN Section	EKACON 107 417 DREW NEKE NEKE DEKE DEKE DEKE DEKE DEKE DEKE
bovine coronavirus poi rah (3933) Human corona 229E poi 1ab (4020) Consensus (4098) avian infectious bronchitis poi 1ab (3989) bovine coronavirus poi 1ab (3989) Human corona 229E poi 1ab (3672) Murine hepatitis poi 1ab (4056) Consensus (4135)	MARCHINA MAR	A TT ROE EYBLAKKN 40 DEDLA O GEBRAA DESS O LERORAVO	LDERS YENAYA LDERKA YDEARA 4150 RELUS REL	DSKNGGVT SCAR	OBCIKOT POID KOT ODO KOT ODO KOT Sectio LEE SYN LEE SIN LEE SIN YKEARIN ON SECTION	EKAMPEKAMPEKAMPEKAMPEKAMPEKAMPEKAMPEKAMP
bovine coronavirus por 1ab (39337) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096) avian infectious bronchitis pol 1ab (3509) bovine coronavirus pol 1ab (3672) Human corona 229E pol 1ab (3672) Murine hepatitis pol 1ab (4056) Consensus (4135) avian infectious bronchitis pol 1ab (3548)	MARCH GMP EN NMARCH NMPSFVI 4135 41 174 82 NAMAGE ITAKSA NIAKSAI	A TUTE ROE SELECTION OF THE PROPERTY OF THE PR	LDERS YENAYA YDEARA 4150 KELDS KELD	DSKNGGVT SCANO SCSAN 4180 ER MTTN DT LTN EQ AAAII ADLALTNM ADLALTNM	O O O O O O O O O O O O O O O O O O O	EKA EKA On 107 417 DRB DKK NRK DKK DKK On 108 421
bovine coronavirus por 1ab (39337) Human corona 229E pol 1ab (3637) Murine hepatitis pol 1ab (4020) Consensus (4096) avian infectious bronchitis pol 1ab (3509) bovine coronavirus pol 1ab (3672) Human corona 229E pol 1ab (3672) Murine hepatitis pol 1ab (4056) Consensus (4135) avian infectious bronchitis pol 1ab (3548)	MARCH GMP EN NMARCH NMPSFVI 4135 41 174 82 NAMAGE ITAKSA NIAKSAI	A TUTE ROE SELECTION OF THE PROPERTY OF THE PR	LDERS YENAYA YDEARA 4150 KELDS KELD	DSKNGGVT SCANO SCSAN 4180 ER MTTN DT LTN EQ AAAII ADLALTNM ADLALTNM	O O O O O O O O O O O O O O O O O O O	EKACON 107 417 DROV NEKK DKKS DKKS DKKS DKKS A21
bovine coronavirus poi rah (3933) Human corona 229E poi 1ab (4020) Consensus (4098) avian infectious bronchitis poi 1ab (3989) bovine coronavirus poi 1ab (3989) Human corona 229E poi 1ab (3672) Murine hepatitis poi 1ab (4056) Consensus (4135)	MAREN GEPTEN NAREN NMPSFVI 4135 41 174 8V 174 8V NIAKSAI NIAKSAI 4174 KVSSII KVSSII KVSSII	AUTONICA CONTROL OF THE CONTROL OF T	A150 A150 A150 A150 A150 A150 A150 A150	DSKNGGVT STAND NGSS SGSAN 4180 SGSAN SGSAN AUDITION ADLALTNM ADLALTNM AUDITION	OBOLKOL POTEKOL OCOLKOL Section INTERPRETARY INTERPRETARY INTERPRETARY OCOLO O	EKACON 107 DREFE DREES DE KES DE KES DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE KES DE LOS DE LOS DE KES DE LOS DEL LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DEL LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DEL LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DEL LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DEL LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DE LOS DEL LOS DE LOS DE LOS DEL LOS DEL LOS DE LOS DELO

***************************************					- Section 109
(4213)	4213	4220	4230	4240	425
avian infectious bronchitis poi 1ab (3587)	FIVCS	NKUTUVIE	PETWVKC	LEGVHOTYST	VVVNIDT
bovine coronavirus pol 1ab (4047)	SLAP	NTLUITVE	DKSVYDOW	VDNVY TTAG	TOLOWA
Human corona 229E pol 1ab (3750)	PATSA	ARTVVVVP	DHDSTYKM	AV DGEVH AC	VANTLOE
Murine hepatitis pol 1ab (4134)	#SLTS	NTITLIVE	DKOVEDOV	VDNVY VT AC	NVNHIGE
Consensus (4213)	PSLSA	NTLTIIVP	DKDVFVQV	VDNVYVTYAC	
					Section 110
(4252)		4260	4270	4280	
avian infectious bronchitis pol 1ab (3626)	ITACE	TELHPTST	GSGLTYCI	SGANIAWPOR	CVNETRNGI
bovine coronavirus pol 1ab (4086)	QDS	THKOLNEI	,SD	DCNWP	LIANRHN
Human corona 229E pol 1ab (3789)	KIND	KNVHLKDV	TK	ENQEI	WPLILTC
bovine coronavirus pol 1ab (4086) Human corona 229E pol 1ab (3789) Murine hepatitis pol 1ab (4173)	QUADS	AVKOLNEJ	DV	NSTWEEN	TAANRHN
Consensus (4252)	QDADO	TNKQLNEI	S	N NWPL	I LNRHN
					 Section 111
(4291)	4291	4300	,431	0	432
avian infectious bronchitis pol 1ab (3665)	NKVD	VLQNNEEL	HGVKTKA	CVAGVDQAH	SVESKEY
bovine coronavirus pol 1ab (4116)	VSAT	TO-MITEL	PAKLKTOV	VNS-GPDOT	NEPTOCY
Human corona 229E pol 1ab (3819)	RVVKI	QWALT	FGKMKVKA	TKG-EGDGG	TSEGNAL
Human corona 229E pol 1ab (3819) Murine hepatitis pol 1ab (4203)	VSTV	IJQ-MILLI	TOKLETOV	VNS-OSDMNO	NTETOCX
Consensus (4291)	vsv v	LQ NNELM	PAKLKTQV	VNS G DA C	CNTPTQCY
					Section 112
(4330)	4330		43	50	436
avian Infectious bronchitis pol 1ab (3704)	TNISC	NSVVA	SSNPNLLV	ASFLNEATNO	TYVDLDD.
bovine coronavirus pol 1ab (4153)	NNSWI	ICKTYY ZII	SDVDGLKY	TKILKDDINI	VVIELDE
bovine coronavirus pol 1ab (4153) Human corona 229E pol 1ab (3855)	UNEGO	RAFMY	TTKECMLY	VEWEHDSS-V	VTVE E
Murine hepatitis pol 1ab (4240)					
Consensus (4330)	NNSGO	GKIVYAII	SD PGLKY	TKILKDDGN	
A-M					 Section 113
(4369)	4369			4390	440
avian infectious bronchitis pol 1ab (3743)					
bovine coronavirus pol 1ab (4192)	CKETI	ODVKGLK1	KTAYEVIG	CNTLARGWY	VETISST .
Human corona 229E ool 1ab (3893)	CREVE	COTPTAPO	KAYA TANKANI	T.NINTERECAL	YTGATO
wurine nepaulis por rab (4279)	Milrot S	CUVKGLKI	TAX TAX VIAG	ENTLASE WE	11111111111111111111111111111111111111
Consensus (4369)	CKFS	QDVKGLK1	KYLYFVKN	CNTLARGWV	LGTISSTV
		<i></i>			Section 114
(4408)	4408		420	4430	444
avian infectious bronchitis pol 1ab (3782)	LOSKO	SHETTEVO	veitsics	FAVLEDADTY	CKYVAAON
bovine coronavirus pol 1ab (4231)	LOAG-	TARLYASI	SSILSLA	SUDPERTY.	DFTQQGG
avian infectious bronchilis pol 1ab (3782 bovine coronavirus pol 1ab (4231 Human corona 229E pol 1ab (3932)	LOAG-	- KQTHEVS	SHLLTHES	EALDPARA	LDAVKQCA
Murine hepatitis pol 1ab (4318)	LOAG.	TATEYASI	SALISTICA	BSMDEKKTYJ	LDYIKQEG
Consensus (4408)					

	<u>-</u>			- Section 115
(4447)	4447		4470	
avian infectious bronchitis pol 1ab (3821)	ELGNOVA'IN	TVHNGSOFAT	SKESPTPHONE	
bovine coronavirus pol 1ab (4269) Human corona 229E pol 1ab (3970) Murine hepatilis pol 1ab (4366) Consensus (4447)	PTANIVKNI	CDHACTEM I	WKPDATTNE	SYCOLOGICAL
Human corona 229E pol 1ab (3970)	PAGNOTHA	TNGSCSCORT	CTIDSNET	MYCTE SOCE
Murine hepatitis pol 1ab (4356)	ZVTNESKINI	COHACTOMAT	DIKPEATUNCE	STEGRATOR
Consensus (4447)	PVGNCVKMI	TDHAGSGMAI	TIKPDATTNOD	SYGGASVCI
				- Section 116
(4486)	4486	4500	4510	
avian infectious bronchitis pol 1ab (3860) bovine coronavirus pol 1ab (4308) Human corona 229E pol 1ab (4009) Murine hepatitis pol 1ab (4398) Consensus (4486)	PERAHTAH:	GSVGNEDGR	OFKESEADTET	TEKREZOFC
bovine coronavirus pol 1ab (4308)	A DEAR VEHI	DVINGLE	KIRCKEVOVEV	GTRIBUSVO
Human corona 229E pol 1ab (4009)	CEAHVAN	TMMET	OYKEKWATU	GTNETTER
Murine hepatitis pol 1ab (4395)	CRERVET	DVASTA	KTROKESTATOT	ETRODICO III
Consensus (4486)	YCRARVEHE	DVDGLC	QLRGKFVQVPI	GIKDPUSEV
				— Section 117
(4525)	4525 4530	4540	4550	
avian infectious bronchitis pol 1ab (3899)	LRNKTOTA	OCKIGYGEOR	DEL BORKGGHO	A 35 E
bovine coronavirus pol 1ab (4343)	LEHDROOV	GEWENGS	VSAPD	TON NO POR
bovine coronavirus pol 1ab (4343) Human corona 229E pol 1ab (4044)	PENTYCKY	GCWLNHGOTO	DRAFI	
				SQFQSKD
Consensus (4525)	LTNDVCQVC	GFWRDGSCSC	VST	SAIQSKD
				- Section 118
(4564)	4564 4570	4580	4590	
avian infectious bronchitis pol 1ab (3938)	KNYGARVEG	SS-ENRLTPL	SCOPPOVKE	STONE NEEDS
bovine coronavirus pol 1ab (4373)	THE SHEET, 27	TS VENRLVPC	ASSESTONO	LEGITO DEV
avian infectious bronchitis pol 1ab (3938) bovine coronavirus pol 1ab (4373) Human corona 229E pol 1ab (4072) Murine benatitis pol 1ab (4482)	N S X Tar Land	SSEARGEPO	CNTDIDYCM	AFOVY ROA
Consensus (4564)	TNFLNRVRG	SSVDARLVPCA	ASGLDTDVOLR	AFDICNKDA
				- Section 119
. (4603)	4603 461	10 4620	4630	
avian infectious bronchitis pol 1ab (3976)	AGMFONLER	MEARTOEEROT		EX Lancor Capture
bovine coronavirus pol 1ab (4412) Human corona 229E pol 1ab (4109) Murine henatitis pol 1ab (4499)	AGIGLHLEV	NUCKFORVDEN	IGDKB	PRVERENT
Human corona 229E pol 1ab (4109)	SFIGKNIKS	NGVPEKNVDKI)	FYTURCIK
The reputate per rap (4499)	MULICULIAN	EXCLUSION FRANCISCO		FEVUERTNT
Consensus (4603)	AGIGLNLKV	NCCREQRVDE	GD KLDA	FFVVKRT L
				- Section 120
(4642)	4642 4	650 .46	60 467	
avian infectious bronchitis pol 1ab (4015) bovine coronavirus pol 1ab (4447)	SNYEHERSC	Tripatale miles	Charles and the Control of the Contr	FVNTCN
11411411 COLOTTA 228E DOL 180 (4 140)	SUMBHEOSM	SOME THE COUNTRIES.	TENTO TO TO THE PROPERTY OF CO.	DITT IL CATTE OF
wurine nepalius por tab (4534)	ENYNKEKEC	THE THER CECAMAL	CE SEE THE PARTY	CODYDUCTOR
Consensus (4642)	SVYNHEKSC	YELLKDC VVA	EHDFFTFDVE	GSRVPNTVR

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(4681)	4681	4690	4700	4719
avian infectious bronchitis pol 1ab (4050)	QRETIKYTM	I PEYAUTH	PEPKULEVLKET	LVIYGCIEDY
bovine coronavirus poi 1ab (4486)	KDLTRYTHI	建工业X为 共民共	FURNICHELCOL	IS I YAGGEOR
Human corona 229E pol 1ab (4179)	ODETRYTH	TLUESTRN.	ROPKINGEVEKE	VLTGCCSTD
Murine hepatitis pol 1ab (4573)	KOTSKETLI	of LdY and H	CLENTESTIKET	TLTYARCEES
Consensus (4681)	QDLTKYTMI	LDLCYALRH	FDRNDCEVLKEI	LVTYACCEDS
				Section 122
(4720)	4720	4730	A740	4758
avian infectious bronchitis pol 1ab (4089)	HPKWFEEN	DIYDPALN	SKYYVMLAKMG	IVERHLLNAI
bovine coronavirus pol 1ab (4525)	1TY	DAXERVEN	ROTATINVYKKE JP	IFNR INSAT
bovine coronavirus pol 1ab (4525) Human corona 229E pol 1ab (4218)	YEM	NEFTER	EDIHRVYAALOK	VVANAMIKCV
Murine hepatitis pol 1ab (4612)	XEOM	DUYEFVEN	POLITINAAKKILLE	IFNETITINTA
Consensus (4720)	Y FEKI	KDWYDPIEN	PDIINVYKKLGP	IVNRALLNAI
				— Section 123
. (4759)	4759	4770	4780	4797
avian infectious bronchitis pol 1ab (4128) bovine coronavirus pol 1ab (4560)	EFCNLMVEN	KINY WE WITH	Sharm nekeyire	SHEOKEABGA
bovine coronavirus pol 1ab (4560)	ETADKLYE	VELL STITE	DNUOTNIKWYTY	DOYNIAADGO
Human corona 229E pol 1ab (4253)	ACCDEMOL	KEVNEVELL	DNODENDNETE	CATVLCPRCM
Human corona 229E pol 1ab (4253) Murine hepatitis pol 1ab (4647)	KHADALAL	ACLULVITE	DIPOLLYCOWN P	GREVKTVPGC
Consensus (4759)	EFAD LVE	KGLVGVLTL	DNQDLNGKFYDE	GDFVKTAPGC
				Section 124
(4798)	4798	A810	4820	4836
avian infectious bronchitis pol 1ab (4167)	AVPVFDTY	POTESMPTEA	OTDATAPERXFE	YD-VHKGYKS
bovine coronavirus pol 1ab (4599) Human corona 229E pol 1ab (4292)	TVALADS	TIME THE	MCHAIDCELLYN	NAYRL
Human corona 229E pol 1ab (4292)	CIPYCIST	TEMPLE VMG	FINCHASECEME	SDIFGODEKT
Murine hepatitis pol 1ab (4686)				
Consensus (4798)	GVPVADSY	YSYMMPMLT	MTHALDSELFVN	
				Section 125
(4837)	4837	,4850	4860	4875
avian infectious bronchitis pol 1ab (4205)	YEALKYDY	TEFROESEC	EXEKYNDOB (H)	NCRDISDOR
bovine coronavirus pol 1ab (4633)	EVEVOIDE	TDYKLEHIN	HALLIAN SMPTH	NTVDCQDDRC
Human corona 229E pol 1ab (4331)	FOLTERIDE	TEHKEVILL	KYNKY (GODYILI	DCVDDHDEMC
Human corona 229E pol 1ab (4331) Murine hepatitis pol 1ab (4720)	EDUVOYDE	POFULLUI	REPRHYSMTYET	NTCECELURG
Consensus (4837)	FDLLQYDF	TOHKLELFN	KYFKHWSQDYHI	NTVDC DDRC
				Section 126
(4876)	4876	4890	4900	4914
avian infectious bronchitis pol 1ab (4244)	LIHCAMSM	difference	TSECNLCRKVE	LEVERIATCE
bovine coronavirus pol 1ab (4672)	THEAUEN	IL SMVLEN	INCLE POVEQUE	ZDEVETVVSIG
avian Infectious bronchitis pol 1ab (4244) bovine coronavirus pol 1ab (4672) Human corona 229E pol 1ab (4370)	THEShirk	TITATTIES	TAFCPUCEKVI	MGVEVVATAG
Murine hepatitis pol 1ab (4759)	THEAMER	TEFSMVIIP!	CECELVEDIE	TIDEREVVSIG
0 (4070)	TTHONNEN	TT DOMUTED	TCFGPLVROIF	IDCUDENTICTO

					- Section 127
(4915)	4915	4920	4930	4940	4953
avian infectious bronchitls pol 1ab (4915) bovine coronavirus pol 1ab (4711) Human corona 229E pol 1ab (4409) Murine hepatitis pol 1ab (4798)	YES	DICATRI	QUINTMSFSKM	GL3Q MOFV	SARATLEGT
Human corona 2005 Life (4/11)	SCH X S	BUGIVME	MINDPHRYRL	SLKDYLLYA	ADTALHVAS
Musing handles and 4409	英語電視	O C D V W	KEUNTHSTRL	TITELLOFV.	PERTITAS
Consensus (4915)	YHYK	ELGVVMN	MDVDTHRYRL	SLKDLLQFV	ADPALHVAS
					- Section 128
(4954)	4954	,4960	4970	4980	4992
avian infectious bronchitis pol 1ab (4322)	SNNI	vinters.	ASVCALTRAL	THUTCHERN	NKERTALIA
bovine coronavirus pol 1ab (4750)	ASAL	YЦЬКС€С	FREATTSOV	KEDTAKEN	ODEVET
Human corona 229E pol 1ab (4448)	SPA	VIK TV	SVAALSTIL	PSCHUYERH	NEFF TO CT.
Murine hepatitis pol 1ab (4837)	ASAS	LADREC	es valles cv	EDPVIEN	GOD TO THE
Human corona 229E pol 1ab (4448) Murine hepatitis pol 1ab (4837) Consensus (4954)	ASAL	VDLRTCC	FSVAAITSGV:	FOTVKPGNE	NODEYDET
*					- Section 129
(4993)	4993	,5000	,5010	5000	
avian infectious bronchitis pol 1ab (4361)	EKA	IFKEG S	TPHENTARPHY	personal very	CONTROL OF VINER
Human corona 229E pol 1ab (4487)	RSQ	FFDLGEE	BTERRY FROM	KEDEKER	
Human corona 229E pol 1ab (4487) Murine hepatitis pol 1ab (4876) Consensus (4993)	LSKO	LKPORS	VDCK (SEFFE)	Sept. Walk NA	LANGE OF THE
Consensus (4993)	LSKG	LLKEGSS	VDLKHFFFTOI	GNAAITDYN	TYVKVNDDT
					Section 130
(5032)	5032	5040	5050	5060	5070
avian infectious bronchitis pol 1ab (4400)	WFD. TO	LLECL	PARTICIPATION OF THE PROPERTY OF THE	CR PER PASO	VEGENAL IN CO.
memericpaule por lab (4815)	50 V 52.3.1	Challed W. Carr	EN MILE PREPARE	COPPEDATION	CONTRACTOR CONTRACTOR CO. CO.
Consensus (5032)	MVDIE	QLLFVL	EVVAKYFEIYE	GGCIPASOV	TUNNYDES
					- Section 131
(5071)	5071	.508	0 .5090)	
avian infectious bronchitis pol 1ab (4439) bovine coronavirus pol 1ab (4867)	AUYRI	Trest	REMS-LET	DOTTELLER	KALUL DUIT IP
bovine coronavirus pol 1ab (4867)	ASYPI	有某口位文章	RIVEALSEEP	UETYAVIT	District Design
isianine riepaulis per rap (4554)	And H. Y. W. H.	BU BURNES OF THE STATE OF	RULE OF THE PARTY OF THE PARTY.	The state of the s	The water than the second second
Consensus (5071)	AGYPE	NKEGKA	RLYYEALSFEE	ODEIFAYTK	RNVI.PTI.T
					Section 132
(5110)	5110	.51	20 51:		
avian infectious bronchitis pol 1ab (4477) bovine coronavirus pol 1ab (4906) Human corona 229F pol 1ab (4604)	MICE	TALBAR	NEWS TO ACTION	TE THE NEW DOOR	0140
bovine coronavirus pol 1ab (4906)	OMET	TAISAR	NATA TO LA STATE	Let be by	2.1000000000000000000000000000000000000
Marke Repailed pur lab (4993)	WITTE O'VE	A JULY SE DE POT	TRAD THE ACTOR	I CHECK OF ALE	to all the state of the
Consensus (5110)	QMNLK	YAISAKI	NRARTVACUST	LSTMTCPOF	HOVETVET
, ,				TO THE GROE	HOWCTRST

				Section 133
(5149)	5149	5160	5170	
avian infectious bronchilis pol 1ab (4516) bovine coronavirus pol 1ab (4945) Human corona 229E pol 1ab (4945) Murine hepatilis pol 1ab (5032) Consensus (5149)	VNTRNASTV	egres fysiod	NH RN IOG	ED PILLMOND
bovine coronavirus poi 1ab (4945)	AAUTIGVEN	ThrukeYaggi	DELIRREPKD	DNEVLNEND
Human corona 229E pol 1ab (4643)	VATINATEM	DESTRUCTION OF THE PROPERTY OF	NEL KNEMADO	DDPKEMGWD
Murine hepatitis pol 1ab (5032)	AATRGVP	Tetrrezgeko	DMSRRLIKD	DSZVIMEND
Consensus (5149)	VATRNVPVV	IGTTKFYGGWD	NMLRRLIKDV	DDPVLMGWD
				- Section 134
(5188)	5188	,5200	5210	5226
avian infectious bronchitis pol 1ab (4555)	Y PRODUMENT	NLENDAASLVI	ARGUTHEGSW	SERIERWYN
bovine coronavirus poi 1ab (4984) Human corona 229E poi 1ab (4682)	The fire of the fire	NILHIVSELVE	ARTHEATTSO	SDREYRLAG
Human corona 229E pol 1ab (4682)	APREDICATE	SMERMLSAME	GSPAVICOTA	SEKFYRESD
wuring nepaulis por rap (5071)	王 A E L A E A E E	NUMBERVSSLVE	ARKIDSCCSH	TORRESPAN
Consensus (5188)	YPKCDRAMP	NILRIVSSLVL	ARKHDSCCS	SDRFYRLAN
				— Section 135
(5227)	5227	5240	5250	5265
avian infectious bronchitis pol 1ab (4594)	FCADY STT	DATEGINUSE	egyseckydt	在某个用户 实际扩张
bovine coronavirus pol 1ab (5023)	SCHOWLSEL	ANCRECALAND	GULSIGODPIOT	ATAMSTENE.
Human corona 229E pol 1ab (4721)	LE METER	MYSH GRAPK	AND THE CHIEF	AYR TSVESS
Murine hepatitis pol 1ab (5110)	RCCCVI SET	INCCCONTARE	eers thank	SEADS SENIO
Consensus (5227)	ECAQVLSEI	VMCGGGYYVKP	GGTSSGDATT	
(5000)	5000			Section 136
(5266) avian infectious bronchitis poi 1ab (4633)	5266	5280	5290	5304
house personaling and 4-5 (5000)	IDATSANA	RLUSVITEDAV	ADMIKSBEAE	YOOV PRV
bovine coronavirus pol 1ab (5062)	COT V. ANVG	ALMSCNGNKIE	DLSIRA JKR	SHWIRSD
Human corona 229E pol 1ab (4760) Murine hepatitis pol 1ab (5149)	I JAVASNIN	CMESWINSSIICN	NINNEKLORO	DEDNCYPENS
Consensus (5266)	COVACAMA	ALLSVNG KIE	DEPTHE	TESNAT VAL
Oonsensus (0200)	COAVSANVC	APPSANG KIE	DPSIKWTÖKK	- Section 137
(5305)	5305 5310	5320	5330	
avian infectious bronchitis poi 1ab (4672) bovine coronavirus poi 1ab (5101) Human corona 229E poi 1ab (4799)	VED 04 24 25	DSZU	5330	5343
hovine coronavirus not 1ah (5101)	MAXING CONTRACTOR	A STATE OF THE STA		THURTHYOO
Human corona 229E pol 1ab (4799)	NUTREBUDD	r cvi otuvka		CHUM VICE
Murine hepatitis pol 1ab (5188)	HVDPALSE	VERTNERPOM	ALC: NO.	SALE PER CE
Consensus (5305)	NVDPAFVSE	FYEFLNKHESM	MILSDOGVVC	MESPERSIN
				- Section 138
(5344)		,5360	5370	5382
avian infectious bronchitis pol 1ab (4711)	LAZDESCEP	EVENNENDER	ANSTONUTE	The selection of the
bovine coronavirus pol 1ab (5140) Human corona 229E pol 1ab (4838)	YTANTSAED	OVENTEN	SESEGEVINA	INNEPARE
Human corona 229E pol 1ab (4838)	YIADTSASK	ATLYZINGZEM	STAKCTOTED	USICTURA
Murine hepatitis pol 1ab (5227)	YTANTSAFO	OVI	SEALGWVETO	TEKEPPER
Consensus (5344)	YIANISAFO	QVLYYQNNVFM	SEAKCWVE D	IEKGPHEFC
· ·		· ·		_

·····					Section 139
(5383)	5383	5390	5400	5/10	540
avian infectious bronchitis pol 1ab (4750)	SOFTE	LVEVOGEPR	Printed and All	CHARLES OF THE PARTY OF	CONTRACTOR SOCIETY
bovine coronavirus pol 1ab (5179) Human corona 229E pol 1ab (4877) Murine hepatitis pol 1ab (5266)	SOUTH	LVKMDGDDV	YERVON	TGEGCE	
Human corona 229E pol 1ab (4877)	SOUTH	QEVDENCKY	mer tean du	THE ACCUMULA	THE TOTAL
Murine hepatitis pol 1ab (5266)	KOMPM	HARMDODEA	DEPT BEST	rg green	DINTERNO
Consensus (5383)	SOHTM	LVKMDGDDV	YLPYPDPSR	ILGAGVFV	DDLLKTDS
The state of the s					Section 140
(5422)	5422	5430	5440		
avian infectious bronchitis pol 1ab (4789) bovine coronavirus pol 1ab (5218) Human compa 2295 pol 1ab (4940)	VAVM	YYDATATES	SPIVEFNE	Mark of the Committee	T. # 10 NOT 672.
bovine coronavirus pol 1ab (5218)	ALL LE	REVSIATE	PRVYDENE	ACK WED W	71.EV.
wurine nepautis por tab (5305)	MINISTRE	PRUSTING	VOLUMENT TO A TO	CASOMIX MODER	100
Consensus (5422)	VLLIE	REVSLAIDA	YPLVYHENP	EYOKVEDV	ALEALER DES
					Section 141
(5461)	5461	5470	5480		F 400
avian infectious bronchitis pol 1ab (4828)	YOU'S	ONNEMDESE	DATE THE PARTY OF	MADO PARVE	enter a
bovine coronavirus pol 1ab (5257) Human corona 229E pol 1ab (4955) Murine henstills pol 1ab (5344)	YNDEG	NOTEDSYSV	TISTEDGOR	TOPESTA	N BOX
Human corona 229E pol 1ab (4955)	NKTIN	EGME EIS FEV	TELDERES	MITPELLA	class price.
Consensus (5461)	YNDLG	NQILDSYSV	ILSTCDGSK	FWDESFYK	NMYLRSTV
					Section 142
(5500)	5500	,5510	5520		5538
avian infectious bronchitis pol 1ab (4867)	Lesce	VESTONSON	THE DAY OF MERCHAN	OF PARECE	ATMEND DESCRIPTION OF
114111411 COTORIA 223E POI 180 (4994)	LUALA	L组织基础保持的第	THE PRESENCE OF THE PARTY	DOM: DOM: MINOR	n Cartiff Den
manne nepanns por rap (5363)	In CASA Ver	AT THE COUNTY	Call to the call of the call	MARKET THE PROPERTY OF	A RESIDENCE AND ASSESSED.
Consensus (5500)	LQSVG	ACVVCSSQT	SLRCGSCIR	KPLLCCKC	CYDHVMAT
(5539) avian infectious bronchitis pol 1ab (4906) bovine coronavirus pol 1ab (5335)	5539	5550	. 556	0	5577
avian infectious bronchitis pol 1ab (4906)	DHENV	SINFITGS	OLCCGEAL	TATIO CIM	STECCHE
numan corona 229E por 1ab (5033)	DEREC	ATTOMATOM	TO STATE WALL AT COLUMN	the second second second	TOTAL PROPERTY.
Munite nepatrils por 1ab (5422)	THE Y-V	LSVSINVEN	SECODVNOT	PRINT THEM	BAYABBILL
Consensus (5539)	DHKYV	LSISPYVCN	SPGCDVNDV	KLYLGGM	SYYCEDHK
					Section 144
(5578)	5578	,559	0 .56		
avian infectious bronchitis pol 1ab (4945)	PKLBI	PEVSNOT	TYRANGAC	ENVOORN	OTA TO HIS
bovine coronavirus pol 1ab (5374)	POYEF	KEVMNLMUE	SLYKOSCTE	PYIDDE	RTASCKUT
avian infectious bronchitis pol 1ab (4945) bovine coronavirus pol 1ab (5374) Human corona 229E pol 1ab (5072) Murine henalitis pol 1ab (5451)	PHLSE	PECSACNUE	GLYKSSALS	MDIDVE	KISTSDWS
Consensus (5578)	PQYSF	PLVSNGMVF	GLYKQSCTG	PYIDDEN	KIASCKWS

		1 10		- Section 145
(5617)	5617	5630	5640	
avian infectious bronchitis pol 1ab (4984) bovine coronavirus pol 1ab (5413) Human corona 229E pol 1ab (5111)	IVEP TES	TR CS DS TO BRIEF	A Safety Blaby Mar. o.	0000
bovine coronavirus pol 1ab (5413)	DVDDVTYA	ECTERARTE	CTO CATTLE A	LO CVE CATE
Human corona 229E pol 1ab (5111)	DIRDYKLA	DAKESTERIA	CRITICERTS ON	CC OVEN TENT
wiunne nepatitis poi 1ab (5500)	EVDDOVIDA	AMCARINE TENER	STREET, STREET	EACY TOME
Consensus (5617)	DVDDYILAI	MECTESLKLFA	AETVKATEEAF	KOSVASATT
			TRILL DURL	- Section 146
(5656)	5656	5670	5680	5694
avian infectious branchitis not tab (5000)	What man the	and the second second second	EFFERENCE AND TEXT OF THE PARTY OF	To district the control of the contr
bovine coronavirus pol 1ab (5452)	ORTVERE			ETTERNOT SEV
bovine coronavirus pol 1ab (5452) Human corona 229E pol 1ab (5150) Murine henatitis pol 1ab (5550)	KETOGPRE	T. T. Sevar	Carlo Carlo	TO THE DESIGNATION OF THE PERS
Murine hepatitis pol 1ab (5539)	DETECTED IN	The state of		DOT TO DOME
Consensus (5656)	RETUSDRE	TISHETCKUK	PRINCHES	THECHNORD
	NOT VODICE!	JI DONGIGRAN.	PPDNKNIVETG	Section 147
(5695)	5695 5700	5710	5720	
avian Infectious bronchitis pol 1ab (5062)	OFFICE	ACHICKON FROM	2720	5733
bovine coronavirus pol 1ab (5491)	V Discovilla	BETHRO - STA	NA I SILATI SVI	A PART OF IN
Human corona 229E nol 1ah (5189)	OWNER WITH	MANY CE DOCKON	Photon from the other miles	The state of the s
Murine hepatitis pol 1ab (5578)		CONTRACTOR OF THE PARTY OF THE	COLUMN THE TAX POST	THE PARTY
Consensus (5695)	VICEFVED	CSELTNC UVV	NATITE WOLF	SA HERRESTOR
	12001 VI DI	COMMING VIII	MALITIKESVG	- Section 148
(5734)	5734 5740	5750	,5760	C770
avian infectious bronchitis not 1ah (5100)	SETTIMEST STATESTICS	CHROTTER	the partered skin of	5772
Human corona 229E pol 1ab (5228)	A PHIDE SHE	AB NEED V COLT V	K D D C D M K C D T	THE PARTY OF THE P
Human corona 229E pol 1ab (5228) Murine hepatitis pol 1ab (5616)	MAST SAPORT	A-100 - 100	FECUVO: NO ME	PANTALINA
Consensus (5734)	VASTISAPTI	WPOR NVTCT	OI VENACUE	E CHRY PRICE
		SALES WIIDI	NIASVISVEE!	- Section 149
(5773)	5773 57	80 579	5800	
avian infectious bronchitis pol 1ab (5139) bovine coronavirus pol 1ab (5567)	A WCKOKRTS	With Displaying	IFA WEEK A LOVE OF	24 6 7 20 6 7 7 7
bovine coronavirus pol 1ab (5567)	HAGMERYCE	Marie Double Charles	Takernteven	
Human corona 229E pol 1ab (5267)	LEGKORTAS	(rfillianded)	COLOTOVVD	
Murine hepatitis pol 1ab (5654)	HIGMERYC	Valgebandski	TANTER	理論は、予論を記
Consensus (5773)	LIGMORYTT	VOGPPGSGKS	AT.ATGT.AVYYC	PADVVPTAC
		· E	. DILLODAVIIC	- Section 150
(5812)	5812 F	820 58	30 584	6050
avian Infectious bronchitis pol 1ab (5178) bovine coronavirus pol 1ab (5606) Human corona 229E pol 1ab (5306)	SHANNOAR	EKAFKETKADI	CTG VEORTT	anced steeler
bovine coronavirus pol 1ab (5606)	SHANDAR	PEATRITUIN	CONTRACTOR	Taken to
Human corona 229E pol 1ab (5306)	THATANASTA	ANVTAYSIO	CTOTTAPAD	E-VOC NO
wurine nepatitis pol 1ab (5693)	SHARVONED	EXAHKELNINI	JOHNA VENKAD	TOWN DIE EXTE
Consensus (5812)	SHAAVDALO	EKAHKFININI	CTRIVPAKURI	ADCASKERT
,,				- DOLUME AL

				Section 151
(5851) 5851	5860	5870	588
avian infectious bronchitis pol 1ab (5217 bovine coronavirus pol 1ab (5645) INDTGKK	TEST THAT	EVSCDALING	ESUPTO VERBE
bovine coronavirus poi 1ab (5645 Human corona 229E poi 1ab (5345) ADTTRK	VIRILIBRIE	EMVTOTVVVL	VERILPRADE
Human corona 229E pol 1ab (5345) HNNSAQ	VESTVEAL	EVNADIVVICE	A HCPV nilevi
Consensus (5851) NOTTRKY	VFSTINALE	EVVTDIVVVDE	V. MLTNYELSVI
				Section 152
(5890) 5890	,5900	5910	5928
avian infectious bronchitis pol 1ab (5256) INGIKI HYC	XVVXVQDPA	TE BY BATELING	-SESPEDVENT
Dovine coronavirus poi 1ab (5684) MARTRAK	HYANT IGU. A	TERVELSK	GTLE RYPET
bovine coronavirus pol 1ab (5684 Human corona 229E pol 1ab (5384 Murine hepatitis pol 1ab (5771) MORISYK	HIVIVEDTO	STATISK	GVME TOVE
Consensus (5890) NARISYK	HYVYIGDPA	QLPAPRVLLSK	GTLEPKYFNVVT
				Section 153
(5929)	5929	5940	5950	5967
avian Infectious bronchitis pol 1ab (5294)	NEMVCVK	THEFTAKE	RESKLINDER	THUSINGSETANN
Human corona 229E pol 1ab (5423	ORMOAIG	MUVE THE L	BE PARTING	ELWENEFUDUR
Consensus (5929)	Krwccre	PDIFLGTCY	RCPKEIVDTVS	ALVYENKLKAKN
				Section 154
(5968)	5968	,5980	5990	
avian infectious bronchitis pol 1ab (5333)	PERRECE	IVIVNNGNS	DVGHESGBAYL	TONEFVEDEVC
bovine coronavirus pol 1ab (5762) Human corona 229E pol 1ab (5462)	ESSSE	HVXXXXG	VTORESSSAVA	O LYLINKELK
Mudos banettis and data (5462)	EARONE	ISLEERG	SVQVDNGSSIN	REDUDVVERETH
Human corona 229E pol 1ab (5462) Murine hepatitis pol 1ab (5849) Consensus (5968)	DINGSIME	KAXXRC	QTTHESSEAVIL	OCTHEESKELK
Consensus (5968)	EASSLCF	KVYYKG	VTHESSSAVNI	100IHLIKKFLK
(0.000)				Section 155
(6007)	6007	6020	6030	6045
avian infectious bronchitis pol 1ab (5372) bovine coronavirus pol 1ab (5798) Human corona 229E pol 1ab (5498)	REKOKRE	AT THE YUA	MIQRAYAMEST	VOIPDSSOOS
Human corona 2005 - 1 d 1 (5/98)	AMPLIPHK	AVELSE TOS	OMFARKAVLCIC	TOTABLACES
Mudpo honetitis and data (5498)	KISTUSK	EVEL SPENIS	QUYVAR LEGG	DOTVESAGESE
Human corona 229E pol 1ab (5498) Murine hepatitis pol 1ab (5885) Consensus (6007)	PHERMEN	WITH JULY S	QNYVAKEVEST	DEPVISAUGER
Consensus (6007)	ANPSWSK	AVETSPYNS	ONI WILL DED TIT OF	TOTTOTTOTTOTT
			ΜΙΙ ΛΗΥΚΑΤΘΤ ζ	
			ON I VAKKATET	Section 156
(6046)	6048	coc	0	Section 156
(6046)	6046	606	0 6070	Section 156 6084
(6046) avian infectious bronchitis pol 1ab (5411) bovine coronavirus pol 1ab (5832)	6046 4 0 Y V J E C	606 VIADSQEAL	0 6070	Section 156 6084
(6046) avian infectious bronchitis pol 1ab (5411) bovine coronavirus pol 1ab (5837) Human corona 229E pol 1ab (5537)	6046 4:1:YV EC 70.Y 1 YS	606 VIADSQEAL O ALTAVSV	0 6070 ITNF ENV. L. 90 EVORTEN L. 20 22	Section 156 6084 CRUTLVVIROR KETICVIENM
(6046) avian infectious bronchitis pol 1ab (5411) bovine coronavirus pol 1ab (5837) Human corona 229E pol 1ab (5537) Murine hepatitis pol 1ab (5924)	6046 LOY IYS LOY IYS	606 VIADSQEAL VIADSQEAS VIADES	0 6070 JTPFE PL TOP LVGETNUS EO PA JANDER VATER	Section 158 6084 REPLEVIER OR EKSTLEVIER OR EKSTLEVIER OR
(6046) avian infectious bronchitis pol 1ab (5411) bovine coronavirus pol 1ab (5837) Human corona 229E pol 1ab (5537) Murine hepatitis pol 1ab (5924)	6046 LOY IYS LOY IYS	606 VIADSQEAL VIADSQEAS VIADES	0 6070 JTPFE PL TOP LVGETNUS EO PA JANDER VATER	Section 158 6084 REPLEVIER OR EKSTLEVIER OR EKSTLEVIER OR
(6046) avian infectious bronchitis pol 1ab (5411) bovine coronavirus pol 1ab (5837) Human corona 229E pol 1ab (5537) Murine hepatitis pol 1ab (5924)	6046 LOY IYS LOY IYS	606 VIADSQEAL VIADSQEAS VIADES	0 6070 JTPFE PL TOP LVGETNUS EO PA JANDER VATER	Section 156 6084 CRITILVVIEROR KETICVIENM

						- Section 1	10/
(6085)	6085	6090	,610)	6110	6	6123
avian Infectious bronchitis pol 1ab (5450)	DELY	SALKE	ELDSETS-		-LOGTGE	HTENRE	de s
DOVING CORONAVIRUS DOI 12D (58/6)	DIFF	ALOFF	PETROKIO	שות שונו עונ	TOP POR NET IS	ATT TO THE PERSON	14 min
wurine nepalitis por 1ab (5963)	OTEF	SLNFT	TETEDRE N	NPR	ROS THREE	Man Cpc	STET?
Consensus (6085)	OLFE	ALNET	LTLDKIN	R	LQCSTNLE	L DCCK	SH.
					HOCSINDE	- Section 1	
(6124)	6124	6130	61	40	6150	-	
avian infectious bronchitic not tab (6402)	Lairy's 676	Me To de un arrive		BOWN DERENE	275 72-20-21-21	Programme and	3162
bovine coronavirus pol 1ab (5915) Human corona 229E pol 1ab (5609)	GYHD	AHEPS	T. ASPIRITORY	D THE COLD	TALVNVEA	GSE	KH
Human corona 229E not 1ah (5609)	DT. DE	Objective in the	To Tober	m	A CHOLON	S-AVIO	N.B.
Murine hepatitis pol 1ab (5999)	222	A W DOO	T WHEN Y	10000	AVOLUNN N	VC41	GE H
Consensus (6124)	CVHD	AUADOL	HAT DDKY	W. C. D. T.	A CUNYAL	Z-WASA	65.K
,	01111	MMESI	DALDOKI	(A2GDT	AACTNAAD	S AVTY	SR
(6163)	6162	6170		180			
avian infectious bronchitis pol 1ab (5521)	#200A	7 35517872	Notes Village of the	GD and a street of	6190	6	201
hovine coronavirus not 1ah (5052)	7	La serie de La	WINVE SCH	MILTR	DESTRING	GUVEEN	ME
bovine coronavirus pol 1ab (5953) Human corona 229E pol 1ab (5646) Murine benefitis pol 1ab (6037)	M 35	CARD LETTER	VIETUDEXCX	+1.4	FR ARES	AMVCED	A
Murine hepatitis pol 1ab (6037)	THE PARTY OF THE P	THE PROPERTY OF	NOMP SH	LECTR	DEAMRHAIL	GULGMI	V.
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(6553)	6553	6560	6570	6580	659
avian infectious bronchitis pol 1ab (5909) bovine coronavirus pol 1ab (6340) Human corona 229E pol 1ab (6031)	FPDSSP	ETHQVDGV	AQ-DLYSLA	TKDCTTKC	TO CAY
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Human corona 229E pol 1ab (6031)	YEDDGS	EVWHDQVN	XXPXR	ATNOTUKER	LOGALI
Murine hepatitis pol 1ab (6424)	MYSDTP	DVYMEGMES	KONDAMBUR	SATOTTRAN	LÉSTAVO
Consensus (6553)	YYSDTP	CVYMDGMDA	KQVDYVPLR	SATCITKON	IGGAVO
				Se	ction 170
(6592)		6600	6610	6620	663
avian infectious bronchitis pol 1ab (5947)	KRUZIOM	AEEVTS	AAVTUSTTE	TNKLNP	ILLIKS
bovine coronavirus pol 1ab (6379)	DOMEE	REYDDOT	TARTICETE	WYKTEDES	JEWN'T
Human corona 229E pol 1ab (6066)	SELANL	RAYVEFT	IFTOATHNI	WYPTERDO	NT TOT
Murine hepatitis pol 1ab (6463)	TENABE	REYLES	TARTEFAR	OWNER OF	HILLIAND
Consensus (6592)	LKHAEE'	YREYLESYN	TATTAGETE	WVYKTFDFY	NLWNTI
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(6631)	6631	.6640	.6650		666
avian Infectious bronchitis pol 1ab (5986)	SIATIL - T	TO STANKE	YKGGHYNAT	ASTMETVIT	Chicagon Com
bovine coronavirus pol 1ab (6418)	4K13	TEN VIVY	TKT HVTCO	agemerati	NUMBER
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Murine hepatitis pol 1ab (6502)	The state of the s			ACCLECAVE	
Consensus (6631)		SLENTVYNT	VNAGHEDG	AGELPCALI	COKVE
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(6670)	6670	6680	.6690		670
avian infectious bronchitis pol 1ab (6023)	IDOGVE	KANTENDONIN	Had syne	Vinceto M. T. D. T. T.	MNDT
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Consensus (6670)	KIONED	VVVFVNNTT	TPTNVAVEL	FAKRSTR H	PELKII
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. (6709)	6709	6720	6730		674
avian infectious bronchitis pol 1ab (6062)					TELET
bovine coronavirus pol 1ab (6494)	BAL MID	CHEBRIT	WADE TENC	NATIONAL MARIN	DAY BY
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Consensus (6709)	KNINID	China and American	VAKECHICE	NUARTH	DI DELL
	MALINED	VINKHVIWL	TAKESFICS		ction 174
(6748)	6748	.6760	.677		678
(6099) avian infectious bronchitis pol 1ab	NCTAG	0/00	(57 mt a fair a fair a)	T HOMO OVER	0/8
hoving coronavirue not tob (6532)	MADMAL	TOUR THUS	S. LARDUAL	LIVE TUC YER	INTAL
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Murine hepatitis pol 1ab (6617)	DIVICT CIT	AWS LUES YE	KILPLETNA	LESATAYKT	GGKS

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Consensus (6787)	PPRADLNG	MVDKVGDSDV	FWFAVRKDGN	DVIFSR DS
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(6826)	6826	.6840	6850	
Quian infectious browning 4				6864
bovine coronavirus pol 1ab (6611)	FRITTIN NOTE	SOCIAL SILE DE	GAFVTLPN	TINENGSSY
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Consensus (6826)	T WOUNDE	SEN PROMING	PESGNEALARG	
	n Apur Pr	CONGN G	L GNDALA	TIFTQSRLL
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(6865) avian infectious bronchliis pol 1ab (6180) bovine coronavirus pol 1ab (6650) Human corona 229E pol 1ab (6312) Murine hepatiits pol 1ab (6734)	6865 6870	6880	,6890	6903
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Consensus (6865)	SSFTPRSDM	EKDFLALDDD	FIOKYGLEDY	APRHTUVCD
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(6904)	6904 6910	6920	6930	
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			SMPATOREA	SYDSSIRSY
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bovine coronavirus pol 1ab (62728) Human corona 229E pol 1ab (6390) Murine hepatitis pol 1ab (6812) Consensus (6943)	TE AT 2 DIG-	PANCETANE	PLUFFIELLE	LKEYGIN
Human corona 220E not tab (6726)	TIDEKS 6-	GSKEVGDVILL	LILEDEVALVE	SENDINCUS-
Murine henotitic not tab (0390)	CIMIATRIDE	SELLACTAMOT	LIL GEVSYLK	EDETVVS-
Concensus (6042)	LIT DENS G	SEKSYCTYIDE	LULIDEV DIVK	SINIKCVS-
Consensus (6943)	FIVDE SG	SSKSVCTVIDL	LLDDFVELLKS	SLNL CVS
				- Section 180
(6982)	6982 69	990 .700	0 7010	7020
avian intectious bronchitis pol 1ab (6295)	KSKYVTVST	PYHSINPUTUE	EDGSTKMCTO	TOZO NIMO
(6982) (avian infectious bronchitis pol 1ab (6295) (bovine coronavirus pol 1ab (6765) (Human corona 229E pol 1ab (6428)	EVNUNV	DEKDEOFULA	NDEKUMBERIDE	ANT
Human corona 229E pol 1ab (6428) - Murine hepatitis pol 1ab (6848) -	KKHEVII	PNKPWRWMT	KONAVATE	T TO KEE
	KVVNLNV	DEKDFORMING	NEWWOOD	THE PERSON NAMED IN
Consensus (6982)	KVVNVNI	DFKDFQFMLWC	NDEKUMERVAL	MANANAN

					Section 181
(7021	7021	7030	7040		705
avian infectious bronchitis pol 1ab (6332)	CHENE	DAY A OUC AW	PCNIPUE	VGITT	S. ILM. K.
bovine coronavirus poi 1ab (6802	Perpusy	DCCXLNSPM	ERVSLWULL	KPVTLI	TOCMMEN
Human corona 229E pol 1ab (6464	CHASIM G	I TORMCL	PCNLY	AGLKUP	SCHMFINE
Murine nepatitis pol 1ab (6886)	PETVETV	LES PL	BRVNEWMER	KPIDEF	TGCMMI
Consensus (7021	PGYSMPV	LYKYQNSPL	ERVNLWNYG	KPITLE	SGIMMNV
					Section 182
(7060)	7060	,7070	,7080		709
avian infectious bronchitis poi 1ab (6371) ETTOELD	ATSKUNIC/	SHAME VMHE	CAGSDE	Greess
bovine coronavirus poi 1ab (6841 Human corona 229E poi 1ab (6503	(P. 10 1 10 1	LNTIDIAV	EVEMBELL T	CHUSER	VEFGSA
Human corona 229E pol 1ab (6503)) Witterer	FNSTLECT	e in course ha	GAGSDY	CLUTTA
Murine hepatitis poi 1ab (6925)	Exprise	LLST MEAN	PANNRULLI	CAPSOK	GUAZGSA
Consensus (7060)	KYTQLCQ	YLSTTTLCV	PHNMRVLHI	GAGSDK	GVAPGSA
					Section 183
(7099)	7099	,7110	.7120		713
avian Infectious bronchitis pol 1ab (6410)	EROWEPE	COLLUSION	VDYCHAHV	SVIST	NKYNTEH
bovine coronavirus poi 1ab (6880	LEOVER	GTT EVENUE L	YPRICESON	TYPEDE	INTERNO
Human corona 229E poi 1ab (6542	KRULLH	VONCTAC	VDY ADE	SVTGDE	ATVYLED
Human corona 229E poi 1ab (6542 Murine hepatitis poi 1ab (6964	TROULEA	CSTILL	NEFERIORIZA	STYCHE	THIERDE
Consensus (7099	LKOWLPA	GTILVDNDV	VPFVSDAVA	SYFCDO	TTLPFDC
•					Section 184
(7138	7138	7150	7160)	717
avian infectious bronchitis pol 1ab (6449)	PER VIET	TONDSKR	KHEGETANN	GNDDV	ILUSSES
67138 avian infectious bronchitis pol 1ab (6449 bovine coronavirus pol 1ab (6919	COTTON	MEDPETRNI	JEYNUSK	ber	PTOUM
Human corona 229E pol 1ab (6581	FOLLES	TOGRIKAL	DGEN DSK		TAINGET
Munne hepatitis pol 1ab (7003	WILLIAM	MIDPLEKNE	GEMNVSK		EFLCHLI
Consensus (7138	FDLIISD	MYDPITKNI	GEYNVSK		TYICHFI
					Section 18
(7177	7177	7190	.72	00	721
avian infectious bronchitis pol 1ab (6488	NHEETENS	FRVKVELT	SKHEVTYDI	AODCAN	WOOMACA A
bovine coronavirus poi 1ab (6954 Human corona 229E poi 1ab (6616	DREALGE	TV/LLIPE	OWNAET YKI	MGYEAR	MIN FORN
Human corona 229E pol 1ab (6616	EKLATE	STATEV	NKKLEET	VORESE	COMPLES
Murine hepatitis pol 1ab (7038	DKLALC	SVEINITE'S	SVNAEU SI	MGKFAE	VIII CIN
Consensus (7177	DKLALGG	SVAIKITEF	SWNAELYDI	MOKFAE	WIMFCIN
					- Section 186
(7216	7216	.723	0 7	240	72
avian infectious bronchitis pol 1ab (6527					
bovine coronavirus pol 1ab (6993) KAJISSEG	FLITTINYLG	KPKVETT	NVMH	MICTOR WENT
Human corona 229E pol 1ab (6655) LTSGIDA	PVVCITYLG	DEAOGPETI	NTTHE	CVERRI
Human corona 229E poi 1ab (6655 Murine hepatitis poi 1ab (7077	NATE	PLTST PUTN	KITRTET	NTM	LLYT PARES
) NASSSEA				

(7255)	7255	7260	7270		Section 187 7293
avian infectious bronchitis pol 1ab (6565)	NYL	TSAYST	VACUDERLKA	TPVVNEKTE	The same of the sa
Murine hepatitis pol 1ab (7114)	TMWN	LSYNGVI GGAXSLI	LLSKENCKHKA	IVVVOEKDS!	INEM LS
Consensus (7255)	TVWN	GSAYSLE	DMAKFPLKLKA	TAVVNLK DO	INDLVLS
		-			Section 188
(7294)	7294	,7300		7319	
avian infectious bronchitis pol 1ab (6604)	DIKC	GHT-T AND	VENTSFTSDSF	CTM SEQ ID	NO: 9905
					NO: 9886
1 10111011 6010110 228	BARS	TEXCHESTAGE!	MERCING PONTER	CEC OF SEC IN	NO: 9914
Murine hepatitis pol 1ab (7153) Consensus (7294)	公共200	AND IL VEL	TRKEVEVGDSI	UNUK SEQ ID	NO: 9887

FIGURE 4B

										Section
		(1)		Marian A. Prins	,10		20			
human coronavirus			MSFT	PGKQ	SSS.	RASS	NRSG	NGTHK-	WADO	SDOVE
	corona NP	' (1)	MSET	PGKQ	SSS.	RASE	INRSG	NG TLK-		SDOSR
vian infectious bronchiti		' (1)						CONTENT A	Transit of	The state of the s
mouse hepatiti		' (1)	MSTV	PGQE	NAGO	RSSS	INRAG	NULLEKI	TTWADE	TERGP
(Consensus	(1)	MSFT	PGKQ:	SSS	RASS	INRSG	NGILK	WADO	SDQAR
										Section
		(40)	40		50		.60)		
human coronavirus	OC43 NP	(36)	VOIR	CRRA	or de	TARSO	OBSG	CNYVP	Variable	SESTION FO
	corona NP	(36)	VOTE	PRA	ó=1.0	TATS	LPSG	ONVVD	Yevrec	77568
vian infectious bronchiti	s virus NP	(16)	TIKL	GPK	PERI	resse.	5-60mm	CONTRACTOR	ALMEOP	LANDE
mouse hepatiti	is virus NP	(40)	NONE	CRRNI	51284	TATT	- Phys	Compat	Yavasc	TUHUU
	Consensus	(40)	VOTE	GRRAC	OPKC) Tare	DSC	CHUUDY	YSWFSG	
					2		2 250	GIVVE		Section :
		(79)	79			0		100		
human coronavirus	OC43 NP	(75)	CKEE	Parrel	LVI C	DE ALO	T D T I	Man Reference	AP HUE C	11 SOUTHER
Bovine (corona NP	(75)	OKER	E DZ					HA HINE	O FIFT A
vian infectious bronchiti	s virus NP	(45)	NAPA	PKF		DNP	LEIV NICH		YT HNIER REQAR	DELTA
mouse hepatiti		(78)	CKEE	OFIA	0.1	FTANC	TONO		YZHNER	
	Consensus	(79)	GKEF	EFARC	SOCI	PTADO	UDAC	FOVCVE	YRHNRR	DATE:
		·· - /					VEAS	POVOIM		Section 4
		(118)	118			130		.140		40
human coronavirus	OC43 NP	(114)	CHOR	OFTER	THE	10 VI 20	ret bar	dia ve	D.P. Company	
Bovine o	corona NP	(114)	CANCOR	OT THE	2 1.73 7541	t with the bear of	Darwin P. Company	13 23 14 15	Towns of the second	Carte in the
vian infectious bronchiti	s virus NP	(82)	CGRK	PMPDA	1	ar is	A STATE	OT MAN	DIDOVE	ASIN
vian infectious bronchiti mouse hepatiti	s virus NP	(117)	chak	OTTO			30	CACACA	CONTRACT.	TO THE PARTY OF
· c	Consensus	(118)	GNOK	OLLO	2.123年で	VVI.CT	CDUA	A D O A C M	STATE SHAPE	WART OF
				22211		11201	GENA	VDÖIGI	PIDGAE	Section 5
		(157)	157			.170		.180		19
human coronavirus	OC43 NP	(153)	SHOW	Tipani	1200	Time of	ida a a a	n'n wan ala	1680	POGYY
Bovine o	corona NP	(153)	E STO	ת תם ת	1000	cutio o r	***			POGYY
lan infectious bronchitis	s virus NP	(121)	AHVE	SESMO	CTE	OKE	NO VICE	DOCEDIO	CDDCNE	and a
mouse hepatitis	s virus NP	(156)	A TON	TRESDA	17	No ele		200	GPDGNE	POCE'Y
· c	consensus	(157)	ADVN	TPADT	TI DD	DDCCF	円のまり	STATES OF		POGFY
		(,					BRIE.	LKEFEG		Section 6
		(196)	100			240		200		
human coronavirus	OC43 NP	(180)	F-5-8-8	D 0 3 D 4	Chic	210	orac an	220	THE REPORT	23
Bovine (corona NP	(1891	PC C	D.C.A.F.	0000	101012	21.43	AGSRS	HANGEN	RTATS
ian infectious bronchitis	s virus ND	(160)	T.C.D	DO CO	DIA S	TOKAS	BHAS	AGSBS	RANSGN	RIPIN
mning density	e virus NP	(100)	Dece	THE LAKE	LAA	DUAAS	V.P.	K	EGS	PGRR
mouse hepatitis	onconcue	(102)	E 2 2 2	PAPP	URS	GSRSQ	n GPI	1NRA	RSISSINO	ROPAS
·	0110011808	(190)	E G S G	KOAPN	ISRS	TSRAS	SRASS	BAGSRS	RANSGN	RTPTS

1000	005 04	^			Section 7
human caranavirus OC42 ND (200	235 24	O New Propagation	250	260	273
human coronavirus OC43 NP (228)	VTPOMAT	OCIA STVI	JAKLGKDAT	KPOOVELHT	KEVROKI
mode nepadie vilus Nr (229	I VKREMA	CILLERY A. L. U.T.	AKTEKDAG	ODVINITION	Crawing a summer of the
Consensus (235	VTPDMAI	QIASLVI	LAKLGKDAT	KPOOVTKOTA	KEVROKT
				22.1121	- Section 8
(274)	274 2	280	290	300	
human coronavirus OC43 NP (267)	TARPROP	CORDINECTO	mer Carron and a	mental Wan	312
avian infectious bronchitis virus NP (227)	PC	WALL DECK		GENUUN	GGEMEKE
mouse henatitis vinus NP (264)	CHARLES TO SEE	FIVERGI	KINDONESE	KTKGKEGNE	DOKMNEE
mouse hepatitis virus NP (268)	HANKER	AND THE PROPERTY OF THE PROPER	SPACCECE	BEENGEN	GSELLKI
Consensus (274)	PNKLKÓK	RSPNKQC	TVQQCFGK	RGPNQ NFO	GGEMLKL
4-1-1					- Section 9
(313)	313	320	,330	340	351
human coronavirus OC43 NP (304)	GTSSPQF	PILABIZ	TTAGATES	SRLEBARVO	N
Bovine corona NP (304)	CTSDPOR	PILARTA	LTACLEES	SRIELEVIE	M
Bovine corona NP (304) avian infectious bronchitis virus NP (205)	GIKDGRV	TAMLNET	SPHACL	BVTPKLOE	DGT. HT. PE
Consensus (313)	GTSDPOF	PILACLA	PTAGAFFF	CSPLETAVIC	NT.
				SCHEED THICAC	Section 10
(352)	352	360	370	000	
human coronavirus OC43 NP (337)		n n n n n n n	AND THE PARTY OF T	380	390
Bovine corona NP (337)		E SECTION OF	ALE PRINTING	UKEDSEESC	PETTMEN
Bovine corona NP (337) avian infectious bronchitis virus NP (301)	PPMMIII	THE REAL PROPERTY.	A RETHANCE	TREDSCISE	EBTIMKY
mouse hengitis virus ND (227)	REILAAL	RESERVE OF E	NINVKICDE	MDGVGRPK	DEVVRPK
mouse hepatitis virus NP (337) Consensus (352)	這樣位	ASSETTING	WEET ON SO	WREDSPIPE	THETTMEN
Odriberiaus (302)	LSGN	DEPQKD	VYELRYNG	IRFDSTLSG	FETIMKV
70041					Section 11
(391)	391	400	410		429
human coronavirus OC43 NP (373)	TMEDIENA	YQQQDGM	MNMSAKPO	ORCHKNGO-	GENEIN
mode richante And Mr (2/2)	TP IN THE LAYOUR	XOKDOGA	DVANCE REDUCT	L TODONOR	KKDEVEN
Consensus (391)	LNENLNA	YOOODGM	MNMSPKPO	SORC KNCO	GENDN
					Section 12
(430)	430	.440	450		400
human coronavirus OC43 NP (409)	T CTELLOW	on trootive	0.50		468
Bovine corona NP (409)	TO THE REAL PROPERTY.	SULFICE	BRETTABL	STERREDER	
					GVVPDGL
Consensus (430)	ISVALPK	SRVQQNK	SRELTAED	SLLKKMDDP	YT
					- Section 13
(469)	469 474			-	- Section 13
human coronavirus OC43 NP (443)	TOW 0 10	SEQ ID N	IO - 9915		
Bovine corona NP (443)	DUISEL	SEQ ID N			
avian Infectious bronchitis virus NP (404)	PLIBET	SEQ ID N			
avian iniectious profitchius virus NP (404)	LGENEL	SEQ ID N			
mouse hepatitis virus NP (449)	EDDSNV		2030		
Consensus (469)	EDTSEI				

FIGURE 4C

					Section 1
	(1)	1 10	20	.30	CV CV
human coronavirus OC43 HE	(1)	W5756007	THO PATORION	YPRATNAVA	STATE OF THE PARTY.
bovine coronavirus HE		METILLPEV	weet role	I a part of a variable	
mouse hepatitis virus HE	747	MARTDAMAPETE	PRATICE SAN DO	Division of the second	
Consensus	(1)	MELLDREL	THAT DOME THE	FNPPTNVVSHL	山口野湖海里 上於
Consensus	(,)	MELLPREI	PASCITESTER	FNPPTNVVSHL	
					- Section 2
	(43)	4350	60	70	84
human coronavirus OC43 HE	(39)	ASSECUTIVE T	NPHYTSLEEDIN	2VIC DESTRES	ENCHSION
bovine coronavirus HE	(39)	FIG PS DELICITIVE OF T	NERHASYMOTH	PARCDSLKINS	CASTONIER
mouse hepatitis virus HE	(43)	Designation of	SQQNERYIDIN	EELEKSCESSA	TALAS DEK
Consensus	(43)	DSRSDCNHIVNI	NP NYSYMDLN	P LCDSGKISS	KAGNSIFR
					- Section 3
	(85)		,100	,110	126
human coronavirus OC43 HE	(81)	SEPPTOR AND AND	ecourt frequ	E PER HASILEN	RSGSADIT
bovine coronavirus HE	(81)	STATE OF THE PERSON	POODIFFEREN	E BUT TO SELLEN	POLONDEN
mouse hepatitis virus HE	(85)	aribid reset	evision average	WEED SUPERIOR	NNOUNE
Consensus	(85)	SFHFTDFYNYTG	EGOOTTFYEGV	NETPYHARKC	SGSNDIW
	,				- Section 4
	(127)	127	140	150	160
human coronavirus OC43 HE bovine coronavirus HE	(123)	Life Various or amount	SALE PROPERTY.	CANADA CARRAMANA	20 m 1 m 1 m
bovine coronavirus HE	(123)	OF KALE VIEW	K NO IN COLUMN	17000	O TTLO
mouse hepatitis virus HE	(127)	MCUKAP THE STATE	OKNER	T CONTROL OF THE STATE OF THE S	CDVCM
		MONKGLFYTOVY			
	(,_,,		KUMAT IKO BIL	VNVLIVINGDA	Section 5
	(169)	169 .18	30 .190	200	210
human coronavirus OC43 HE	(165)	CS	AYTAPOANSCE	TALKV TABLYS	OCCURVATE
bovine coronavirus HE	(165)	BSCVICTOR	AVINEENSPO	WAR THEY	
mouse hepatitis virus HE	(169)	TANGUTTANA	PERFUSED	TOTAL PROPERTY	NAME OF THE OWNER, OF THE OWNER, OF THE OWNER, OF THE OWNER, OF THE OWNER, OWNER, OWNER, OWNER, OWNER, OWNER,
Consensus	(169)	GS LVINNE	AYTAKEAN GO	YYYKVEADFYL	SCCDEALA
	(,		miimkoma oc	TITROBACTIO	Section 6
	(211)	211 220	230	240	252
human compavirus OC43 HF	ໄຂດຊາ	WHAT SHOULD AND	market deplement of	Styles of the star	COLUMN TO THE
hoving coronavirue HE	(203)		TO THE RESERVE		
bovine coronavirus HE mouse hepatitis virus HE	(203)	ALC: NO.			
Concording	(211)	PLCIPNGKFLSN	アロリの北部が設定を開	聯人而知為多等的時中觀	Mile Distriction
Consensus	(211)	PLCIFNGKFLSN	TKIIDUSQIIE	NKDIGATAGIN	STETITIES Section 7
	(253)	253 260	270	280	294
human coronavirus OC43 HE bovine coronavirus HE	(245)	PERMIT TOWN	OU VINTER OF	Tara tara tara	LAND FOR DA
mouse hepatitis virus HE	(253)	T. DE MC TANK A WILL D			
		FDLNCHYLVLPS			

Section 1

49/193

FIGURE 4C (contd.)

										- Section 8
(295	295	300		,31	0			90		336
human coronavirus OC43 HE (287) 為時	HOSEWNI	JALOS	De Car	V.Est	c = c	22 C	PERS	TINT	CAYDEN
bovine coronavirus HE (287) 📖	DETWIN	IAR CE	E PM	rav	Act	Prin	YFRNS	THY	LUYLIN
bovine coronavirus HE (287 mouse hepatitis virus HE (295	(編集)	LUL VIHS	NE	工艺/0	LAI	300	LETU	TORKT	SDA	NEWPSH
Consensus (295) QV	VDSRWN	VAROS	DNM	TAV	ACC	PPYC	YFRNS	TTNY	VGVYDIN
										Section 9
	337			350			360			378
human coronavirus OC43 HE (329 bovine coronavirus HE (329) 報題	dag en s	LESQ'E	LYN	S PS	F 5 (OWVE	RYDN	SSVT	PLYPAIN
bovine coronavirus HE (329)	DAS LUSI	LSAL	IX. D	SP.	ESC	OSVE	RIDNY	SUV	PLYSTER
mouse hepatitis virus HE (337) 解質	DATE SEE	LAGL	M'M	vs.	LA	COAT	VENTE	SSI	POXPAGH
Consensus (337) HG	DAGFTS	ILSGL	LYN:	SPC	FSC	QGVF	RYDNV	SSVW	PLYPYGR
										Section 10
	379		390			A	00		410	420
human coronavirus OC43 HE (371) 作品	LINDIN	NPDL	LCV	LOT	地型	Like	TIRGV	A.V.	VLLL
bovine coronavirus HE (371 mouse hepatitis virus HE (379) 翻	DAYD IN	PEDV	ICV	y by	1027	i i i i i i i i i i i i i i i i i i i	TLLEV	TELL	LYVLLL
mouse hepatitis virus HE (379) 際認	DENI VI	P-MA鬻	AC W		SEE.		VERGI	TANKE !	FMF
Consensus (379) CP	TAADIN	PDPB	ICA.	YDE	PP	/ILLG	ILLGV	AVII	IAAFFFĀ
										Section 11
) 421	***************************************	432							
human coronavirus OC43 HE (413				SEQ		NO:	9916			
bovine coronavirus HE (413				SEQ		NO:	9888			
mouse hepatitis virus HE (420				SEQ	TD	NO:	9899			
Consensus (421) FM	VDNGTR1	LHDA							

FIGURE 4D

				Occupi i
•	(1) 1	10	20	39
bovine coronavirus Sm		AYFADTVWYV		LVIIVVVAFEA
avian infectious bronchitis virus Sm	(1) MN五上版	KSTEBNGSEL	PALYLIVGELA	YLLGRALQATVQ
mouse hepatitis virus Sm	(1)MF190	LETTOTVWYV	SOIIFLFAVCI	MVTIIVVERIA
Consensus	(1) M M N	FL DTVWYV	COLIFIVALCE	LVII IVVAFLA
				Section 2
	(40) 40	.50	.60	78
bovine coronavirus Sm	(38) TEKLC	TOLCGMENTL	V LSPSTYV FNR	GROFYERYN-DVK
avian infectious bronchitis virus Sm	(40) AADAE	CLFWYTWVVI	PGAKGTAFVYK	YTYGRKLNIPELE
mouse hepatitis virus Sm	(36) SIKLO	TOTOGLENTE	VICEPSTYTVINE	SKOLYKYÝNEEMR
Consensus		IOLCGLCNTL		
	(10) 0 1120	TODOGECKI I	ADDEDITION K	Section 3
	(79) 79	90	•	108
bovine coronavirus Sm	(76) PPVLD			* 1,1100.000000
avian infectious bronchitis virus Sm			NPANFQDAQRD	SEQ ID NO: 9889 KLYS SEO ID NO: 9907
mouse hepatitis virus Sm	(75) LELLE		NPANEQUAQRU	SEQ ID NO: 9900
Consensus	(79) PILD	white		DEQ ID NO: 9300

FIGURE 4E

						Section 1
	(1)	1	,10	20	30	4
human coronavirus OC43 M	(1)	-MSSKTTP	APVYINT	ABEALKELKEW	FSECTI	
bovine coronavirus M	(1)	-MSSVTTP	APVYTWT	ADEALKELKEWI	Irsugia	LLF.ITT.
avian infectious bronchitis virus M	(1)	MSN	ANCTLD	CEOSVELFERY	LFITAF	LEGIT
mouse hepatitis virus M	a	MISTITOAP	OPVYOUT	ADBAIRFIRZW	FSLGTD	if virt
Consensus	(1)			ADEAIKFLKEW		
00.1007.000	.,,					Section 2
	(41)	41	50	.60	70	8
human coronavirus OC43 M	(40)	Carrier and Carrier	a distractive start	ANTELWIMAST:		NEW AT
bovine coronavirus M	(40)	FEMMON	MARKA VAT	PATTE STM PATE	PINTE	N WY AIT
avian infectious bronchitis virus M	(36)		S D T T O T M	THE WALLEY OF THE BOOK OF THE	NEDARGALT	SPP
mouse hepatitis virus M	(41)		E WHITE TAKE	BOYLOW DATE	e Wilchie	NOW AT
Consensus	(41)	TOFCVTCP	CMETTALL	KMIILWLMWPL	TITLTE	NCVYAL
Consensus	(+1)	DOEGIISK	SHEVIVI	KHILDWIMELD		Section 3
	(81)	£1	90	100	110	12
human coronavirus OC43 M	(80)	STATE OF THE STATE OF	White Take	IMWLV#EVNSI	STORES OF B	a Pagaria
bovine coronavirus M	(80)			TMOTOSPANIST	e i e r e m	SWEAK FILE
avian infectious bronchitis virus M	(76)	C CHAIN A W	TIT POST STATE	IMWEGYWIQ I USFEGYWIQ I	PKEG	OWNER PRO
mouse hepatitis virus M	(04)	NAME OF THE PARTY	THE TOTAL OF	INVINCENNE I		Carle by
Consensus	(81)	MIZZICEGI	VPTTVAT	IMWIVYFVNSI	RIFIRTG	SWWSFN
Collections	(01)	MAIDGESI	V L L L V 113			_ Section 4
	(121)	121	.130	.140	.150	16
human coronavirus OC43 M bovine coronavirus M	(120)	ETSNEMCT	DMKCTMY	VERLIBOYETL	TVILTEG	HLYIQG
hovine coronavirus M	(120)	RIVINAME	DMKGRM	VRPID ODYHTE	TVTTTRE	HEYMOG
avian infectious bronchitis virus M	(116)	MANAVGS	LISNGO	CNFATESVPMV	LSPIIKN	GVLYCE
mouse hepatitis virus M	(121)	FPINING!	DMKGTV	VREILSDYHTL	TAPITEG	HEYMOG
Consensus	(121)	ETNNLMCI	DMKGTM	VRPILEDYHTL	TVTIIRG	HLYMQG
	(,					- Section 5
	(161)	161	.170	,180	.190	20
human coronavirus OC43 M bovine coronavirus M	(160)	KLCTCYSW	ADTHAY	NTVAKVTHLCTY	KRGELDE	ISDTSG
hovine coronavirus M	(160)	KLCTCYSI	SDUPAY	VIWAKVSHLLIT	KRGELDI	TGDTSG
avian infectious bronchitis virus M	(156)	OWLAKCE	DHAPKD	TENCPPDERNI	R	N
mouse hepatitis virus M	(161)	KTGTGEST	SDECAY	VIVARVSHICT	KRAFLDE	VDGVSC
Consensus	(161)	KLGTGYSI	SDLPAY	VTVAKVSHLCTY	KRGFLDE	KI DISG
						- Section (
	(201)	201	210	220	231	
	ionn	AVVESKI	MIYRLE	STOKESEMDTAT	LENNÍ S	EQ ID NO: 9
human coronavirus OC43 M						EO ID NO: 9
human coronavirus OC43 M bovine coronavirus M	(200	AUYVKSKI	LGMYRLE	STOKESGMDTAL	ERNNI "	
hovine coronavirus M	(200	AUYVKSKI	LGMYRLE	STOKESGMDTAL	ERNNI S	EQ ID NO: 9
human coronavirus OC43 M bovine coronavirus M avian infectious bronchilis virus M mouse hepatitis virus M	(200) (185	QKYTGDQ	CHYRLE CHKKRF	STOKESGMDTAL AT	ERNNI S	

FIGURE 4F

- Section										
4	0	,30		20	78 Cartes 1 7 - 12	.10		1	(1)	
PPIST	DTGP	-INDKI	TSDN	LKCI	fávig	LPTA	LUIS	METIT	(1)	human coronavirus OC43 S
									(1)	avianinfectiousbronchitisvirusS
PSTST	DTGV	-UNDVI	TTVS	же	FAVIG	LPTA	LUTS	MFL	(1)	bovine coronavirus S
PSIST	NVSA	NSNGAL	IQLV	IFRCI	LGYIG	LPSC	FILE	MLF	(1)	mouse hepatitis virus S
PSIST	DTG	IND I	TSL	LKC	FAVIG	LPTA	PPIS	MFL	(1)	Consensus
- Section										
8	0	.70		,60		50		41	(41)	
TYRNM	TSGS	NGYYP	TLF	CTINE!	V.L DRV	GTYY	TNGI	TVD	(40)	human coronavirus OC43 S
CAULVI	CALL	T. VPT. T. C	TO DAME	-MIT. V.C					(1)	avianinfectiousbronchitisvirusS
TYRNM	TSGS	ŇĠŸ¥₽Ţ	LLI	LNT	VLDRV	GTYY	TNGI	TVD	(40)	 bovine coronavirus S
KERNL	VDGS	TGYYPV	11 15 15 11	LINA	VLDRV	GTYX	2001	LVE	(41)	mouse hepatitis virus S
TYRNM	TSGS	NGYYPT	TLLI	LNT	VLDRV	GTYY	TNGI	TVD	(41)	Consensus
- Section										
12		,11		,100		.90			(81)	
KDRVM	エドハゴ	ARVKNI	NUTE	DEI	KEREL	RLWH	VIII.	LKG:	(80)	human coronavirus OC43 S
NNAGS	SSEE	AVMNI	GNA	HLO	RUSG	QSAL	VYY	SSS	(23)	avianinfectiousbronchitisvirusS
KDGVM	TKVI	AKTEM	NUTI	SDET	KEFEL	TLWE	LILLS	DKG	(80)	bovine coronavirus S
TPSGA	LKTS	AK QNI	DULL	QEN	QPSYL	LSWI	NSV	TTS	(81)	mouse hepatitis virus S
	TKVI	AKVKN	NGI	SDFNI	KPPFL	SWF	VLLS	LKG	(81)	Consensus
 Section 										
16		,15		,140	SELECTION OF	,130			(121)	
KLOGL	DGD	INSTO	OPRI	SVVV	VNTSY	GSTE	ATT	SEF	(120)	human coronavirus OC43 S
	s-		TAP-	STAM	V V NA S	HGGR	VGI	SGC	(63)	avianinfectiousbronchitisvirus\$
		н∰т								bovine coronavirus S
	YNG-	!			GYTSY					mouse hepatitis virus S
KLQGL:	GN	T	QP	5 VVV (VNTSY	GSTF	AIT	SEF	(121)	Consensus
 Section 										
20	90			,180		,170	and tracerd		(161)	
		IRKETWI								human coronavirus OC43 S
CPLTG	KHGG	VTHCY	TTVI	YFSD	CTAHE	SBCE	MAW		(88)	avianinfectiousbronchitisvirusS
GWVSC.	SHM DI	RVELWI	G-NC	NPNL	PNII.	MCEA	COX	ET	(156)	bovine coronavirus S
		(LIGFW)								mouse hepatitis virus S
	IH DI	RIELWI	G N	NPNL	PNTIC	MCEY	CQY	EIS	(161)	Consensus
Section							<u> </u>			
24	30			220		210	and a second		(201)	
TOURSET	DIGV	YAYFT	GGT	FYQE	YLYEH	VNAT	T'T'Y	YKR	(199)	human coronavirus OC43 S
V.I. P. L					MCOLE	TORMI	T. TRY	TOO	(126)	avianinfectiousbronchitisvirusS
CVNN	RSFC	KYPTF	VSV	XMTT.	maxmr	DAM	11 -1007 TA M		(120)	
CVNN	DTGV	YAYET	GGT	FYQE	YLYTH	VNAE	FTY	YKR	(195)	bovine coronavirus S
CVNNI VTKF ATTFT	DTGV VDKPS	YAYET YAYYA YAYYA	GGT IGGT	FYQE FYQH	AFYFH	VNAE	FTY)	YKR	(195) (189)	

				·		Section 7
	(241)	241	250	260	270	280
human coronavirus OC43 S	(239)	NVYLGM	AISHYYVM	PUTCNSK	TITEAMALE	LTSRO
avianinfectiousbronchifisvirusS	(166)	SYYLNG	DEVYTSNE	TIDVTSAGVYE	KAGGPITYK	VMREVE
bovine coronavirus S	(235)	NYTLGT	VLSHYYVM	Blechsp	TTLEYNVER	LTSKO
mouse hepatitis virus S	(229)	SVYEGD	ILTOYYVL	PFICNPTAGET	FAPRYNVIP	LVKRQ
Consensus			ILSHYYVM	PLTCN A S	LTLEYWVTP	LTSRQ
						- Section
	(281)	281	290	300	310	32
human coronavirus OC43 S	(275)	LAFNE	DETTENAE	DOMEDEMSELF	CKTOSIAPP	TSVYE
avianinfectiousbronchitisvirusS	(206)	ALAYEV	NETAODVI	LCDGSPRGLE	COYNTGNES	DOFEP
bovine coronavirus S	(271)	TITARNO	DOVTENAV	DEKSDEMSET	KTLSIAPS	TEVYE.
mouse hepatitis virus S	(269)	TFNENC	KAVITSAV	DEASSYTSET	KIOSMLES	TOVYE
Consensus	(281)	LLAFNO	DGVIENAV	DC SSFMSEIR	CKTOSIAPS	TGVYE
	\ ,					- Section
	. (321)	321	. 330	340	350	36
human coronavirus OC43 S	(315)	NCYTO	PIADAYRE	KPNLPNCNIE	WINDKSVES	PUNNE
avianinfectiousbronchltisvirusS	(246)	TNSSLT	KOKFEVYE	E1	SUNTTOTLH	NFIEH
bovine coronavirus S	(311)	NEYTV	RIADVYRA	IPNUPDENT	WINDKSVES	PLNUE
mouse hepatitis virus S	(309)	SGYTVE	PVGVVYRI	VANLPACNILI	EWDTARSVES	PLNWE
Consensus				IPNLP CNIE		
	(,					Section 1
	(361)	361	,370	380	390	40
human coronavirus OC43 S	(355)	KTESNO	TENMS SALE	isficadstec	NNIDAAKING	MCFSS
avianinfectiousbronchitisvirusS	(277)	ENGANI	NP	SGVUNIOTYO	TKTAOSGYMN	PNESF
boyine coronavirus S	(351)	KITSM	NEWNSSE	SPUDAYSE C RYVCAESLEC	VNIDAAKI (C	MCFSS
mouse hepatitis virus S	(349)	KITO	MENISSLI	RYVEAESLEC	NNIDASKVYC	RCTGS
Consensus	(361)	KTFSNO	NENMSSIA	ACETOAD CETC		
				19 ET GWD 2 ET C	NNIDAAKIYG	MCFSS
				43FIQADSFIC	NNIDAAKIYG	MCFSS Section 1
	(401)	401	410	420	430	Section 1
human coronavirus OC43 S	(395)	401	410	420 Lelenievio	430 SERVRIDTTA	Section 1 44 TROD
human coronavirus OC43 S aylaninfectiousbronchitisvirusS	(395)	401 TIDEF	410 MENGREVI KESNEMYGS	420 DEGLENLEYLO SYHPSCKERLE	A30 SERVEL DTTA TIENGEWENE	Section 44 TREOL
	(395) (311) (391)	401 TIDEF SSFVT	410 Q PNGPKVI KESNEMYGS XI PNGRKVI	420 offolganeyro synpsckerie obougangyld	430 SENYRI DTTA TI ONG WENS SENYRI DTTI	Section 44 TREOL LSVSI TREOL
avianinfectiousbronchitisvirusS	(395) (311) (391) (389)	401 TIBER SSFVY TIBER SVDKR	410 Q PRGREVI KESNEMYGS ALPNGREVI AV PRSKOVI	420 DIQUENTEYLO SYHPSCKERLE DEOLEMISTO DADLENSEFLO	A30 SENYRIDTTY DI ANGEWENS STNYRIDTE TANYRIDTAY	Section 44 TREOL LSVSI TEGOL TUCOL
avianinfectiousbronchitisvirusS bovine coronavirus S	(395) (311) (391) (389)	401 TIBER SSFVY TIBER SVDKR	410 Q PRGREVI KESNEMYGS ALPNGREVI AV PRSKOVI	420 DEGLENLEYLO SYHPSCKERLE	A30 SENYRIDTTY DI ANGEWENS STNYRIDTE TANYRIDTAY	Section 44 Tigot Living Tigot Tigot Tigot
avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S	(395) (311) (391) (389)	401 TIBER SSFVY TIBER SVDKR	410 Q PRGREVI KESNEMYGS ALPNGREVI AV PRSKOVI	420 DIQUENTEYLO SYHPSCKERLE DEOLEMISTO DADLENSEFLO	A30 SENYRIDTTY DI ANGEWENS STNYRIDTE TANYRIDTAY	Section 4 THEOT LEVEL THEOT THEOT
avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S	(395) (311) (391) (389) (401)	401 TIDEF SSFVX TIDEF SVDKF SIDEF	410 ALENGRKVI KESNEMYGS ALENGRKVI NYERSKOVI ALENGRKVI	420 DIOLGNESTLO SYHPSCKFRES DIOLGNESTLO DEGLESSET, O DLOLGNESTLO A60	430 SFRYBIDTTF TIANGEWFNE STAYRIDTTF TANGKLOTAF SFNYRIDTTF	Section 4 (T1COI LEVSI TEGOI ATSCOI Section 4
avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S	(395) (311) (391) (389) (401)	401 TIDEF SSFVX TIDEF SVDKF SIDEF	410 ALENGRKVI KESNEMYGS ALENGRKVI NYERSKOVI ALENGRKVI	420 DIGIGNIGYLO SYHPSCKFRIE DEQUGNEGYLO DEQUGNEGYLO DLQLGNLGYLQ	430 SFRYBIDTTF TIANGEWFNE STAYRIDTTF TANGKLOTAF SFNYRIDTTF	Section 4 (T1COI LEVSI TEGOI ATSCOI Section 4
avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(395) (311) (391) (389) (401) (441) (435)	401 ELDEF SSFVX TIDEF SVDEF SIDEF 441	410 YENGEKYI KESHEMYGS ATENGEKYI AYENGEKYI ATENGEKYI ATENGEKYI ATENGEKYI ATENGEKYI ATENGEKYI	A20 DEGENEGALO STAPSCKERE DEGENEGALO DEGENEGALO DEGENEGALO A60 VESTWIKKERGE	A30 SERVELDTTE TIANGENERS SERVELDTA TARVELDTA SERVERDTA A70 IEDSVEKER	Section 4 THEOLE LEVEL THEOLE THEOLE ATSCOL Section 4 AGVLI
avianinfectiousbronchilisvirusS bovine coronavirus S mouse hepatilis virus S Consensus human coronavirus OC43 S avianinfectiousbronchilisvirusS	(395) (311) (391) (389) (401) (441) (435)	401 ELDEF SSFVX TIDEF SVDEF SIDEF 441	410 YENGEKYI KESHEMYGS ATENGEKYI AYENGEKYI ATENGEKYI ATENGEKYI ATENGEKYI ATENGEKYI ATENGEKYI	A20 DEGENEGALO STAPSCKERE DEGENEGALO DEGENEGALO DEGENEGALO A60 VESTWIKKERGE	A30 SERVELDTTE TIANGENERS SERVELDTA TARVELDTA SERVERDTA A70 IEDSVEKER	Section 4 THEOLE LEVEL THEOLE THEOLE ATSCOL Section 4 AGVLI
avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(395) (311) (391) (389) (401) (441) (435) (351)	401 PLDEP SSFVM TIDEP SVDEF SIDEP 441 EGP	A10 TOPICE KYTER KESNEMYGS AIFNGEKYT TYERSKOV ASO ANYSVSKE	420 DIOLGNESTLO SYHPSCKFRES DIOLGNESTLO DEGLESSET, O DLOLGNESTLO A60	A30 SERVALDITA DI DI DI DI DI DI DI DI DI DI DI DI DI D	Section 4 THE OLI LEVEL THE OLI THE OLI THE COLI ATSCOL Section 4 AGVLI G AGVFI

						Ser	ction 13
	(481)		490	500		10	520
human coronavirus OC43 S	(475)	HOVVYAC	HGFKAPF	NECRE	NG-SCYGSC	ž	GKNN
avianinfectiousbronchitisvirusS	(357)	GCK	SVEKGR	TCGYAYS	YGGPSLCKG	VYSG	
bovine coronavirus S	(471)	HDVVYAD	HCFKASI	MEGECKE	DGSLCVGNG	PGIDAG	YKTS
mouse hepatitis virus S	(463)	HDVVYAS	QUETVRS	SYPPCAO	PDIVSPCTT	OTK	E
Consensus	(481)	HDVVYAC	HCFKARS	NECPCKL	G LSVGSG	P	ĸ
						Ser	ction 1
	(521)	521	530	.540) 5	50	56
 human coronavirus OC43 S 	(509)	GIGICE	CHNYLTO	DN	-LCTPOPIT	FEGT	YKCI
avianinfectiousbronchitisvirusS	(388)						ASSESSED NO.
bovine coronavirus S	(511)	GIGTCPA	GTNYLTO	HNAAQCD	CLUPPPP	SKATGP	YKC
mouse hepatitis virus S	(497)					MP788	· HEATH-COM.
Consensus	(521)	GIGTCPA	GTNYLTC	: N	LCTPDPIT	TG	YKCI
							ction 1
	(561)	561	570	580) 5	90	60
human coronavirus OC43 S	(541)	OTKSLVC	ICERTS	LAVKSDY		OTCREC	
avianinfectiousbronchitisvirusS				VYVTKS		The state of the s	
bovine coronavirus S	(551)	OTRYEVE	LEEHUS	LATKSON	CO	CTCORO	A STATE
mouse hepatitis virus S	(497)	-KSAFVA	VEDHEE	TIGVILEDIN	CHADPHKG	CICAN	SPT
Consensus	(561)	QTKALVO	GIGEHCS	LAVKSDH	CG GN		AFL
			····				ction 1
•	(601)	601	610	.620) 6	30	64
human coronavirus OC43 S	(577)	WSADSCI	OGDKEN	FANFITH	DVNSGLTCS	TOTAL L	North Control
avianinfectiousbronchltisvirusS	(407)	25000 - 400,000 - 2		SR	-JOTATEPE	VITTINE	VNN
bovine coronavirus S	(587)	WSVDSCI	OCDRON	FAN ET TH	DVNSGTTCS	Individue	Al mins
mouse hepatitis virus S	(536)	WSHDTGI	VNDRCO	FANILLE	GINSGPPCS	TIDIL PIT P	MTE
Consensus	(601)	WS DSCI	QGDRCNI	FANFILH	DINSGTTCS	TDLOKA	NTD
						Se	ction 1
	(641)	641	.650	660) 6	70	68
human coronavirus OC43 S	(617)	TLGVCUN	PERGLI	COUTEVE	WNAIWYNS	ONTHE	ENG
avlaninfectiousbronchitisvirusS	(427)	TENTONE	NITTER	EAGETTN	TDBAVS	YNY	ACT.
bovine coronavirus S	(627)	TLGVC	DLECT	COCTEVE	VNATYYNS	Cont. Cy.	SNC
mouse hepatitis virus S	(576)	VIGICIA	CYDDYCI	GOGVEKE	VNATYYNSÜ MKADYYNSÜ	OTLINE	VNG
Consensus	(641)	ILGVCVN	YDLYGI	GOGIFVE	VNATYYNSW	ONLTY	SNG
							ction 1
	(681)	681	690	.700	. 7	10	72
human coronavirus OC43 S					RVSAAFHAN		
avianinfectiousbronchitisvirusS	(465)	TLDTSGS	TOTEVVO	GPYGINY	YKVNPCEDV	NOVERN	CCC
bovine coronavirus S	(667)	TYPERO	TTNETE	MERCVEC	RYSAAFHAN	100 0 0 0 V	1000
mouse hepatitis virus S					RVSAAFHKI		
Consensus					RVSAAFHAN		

	(704)	701	730	740	750	ection 19 760
human coronavirus OC43 S	(721)			QLQPINYFDSYLG		
avianinfectiousbronchitisvirusS						
bovine coronavirus S	(202)	TAGILISE	WHICH	OLDENOFYIKITN OLDEINYFOSYLG	CURRERROL	CCUTTO
mouse hepatitis virus S				EENPLNYFDSYLG		
Consensus				QLQPINYFDSYLG		
Consensus	(/21)	INCNIAL	INSUSK	.Žnčiiuiinoiin		Section 20
	(761)	701	.770	780	.790	800
human coronavirus OC43 S				YSKNERSEGALTI		
avianinfectiousbronchitisvirusS	(545)	Will DVII CV	D TO TO	PDG		LALKOT
hovine coronavirus S				YST KRRS KRS LTT		
mouse hepatitis virus S				YSKSKRADBSVST		
Consensus				YSK RRSRRSITI		
Conscisus	(101)	TCDHIVGC	761645	TOR KNORNOTTA		Section 21
	(801)	801	810	820	830	840
human coronavirus OC43 S	(777)	VNDSTEP		TOTESETTIONNY		
avlaninfectiousbronchitisvirusS				MLSSNSTNLTVTI		
bovine coronavirus S	(787)	VNDSVEP	COLUM	TOTASEUTIGNME	OF LOUSS P	CONTRD
mouse hepatitis virus S				модетитттенн		
Consensus	(801)	VNDSLEP	GGL YE	IQIPSEFTIGNME	EFIOTSSPI	KVTIDO
	(,					Section 2
	(841)	841	.850	860	870	88
human coronavirus OC43 S	(817)	AAEUCED		OLVEYUSECONIA	ALLTEVNE	LLOTEC
avianinfectiousbronchitisvirusS				CLEOONEDVEDVII		
bovine coronavirus S	(827)	SAEVICE	YAACK	SOLVEYCSFOULT	PATLTENE	LLDTT(
mouse hepatitis virus S	(776)	AAFVCEDI	NTACRO	OLVE 45 SECVEN	ATLNE NN	ььбим
Consensus				QLVEYGSFCDNIN		
						Section 2
	(881)	881	.890	,900	910	92
human coronavirus OC43 S	(857)	LOVANSL	MNGVEI	erktrochnenni	DINFSPVL	GCLGS
avianinfectiousbronchitisvirusS	(651)	INFYSST:	KP	AGFNTPXLS的证:	STGE NIST	LLTNP
bovine coronavirus S				CSTKLKDGVNENVI		
mouse hepatitis virus S				CSSRIPDGTSGPT!		
Consensus	(881)	LQVASSL	MNGVTI	LSTKLKDGVNFNV		
						Section 2
	(921)		930	940	,950	96
human coronavirus OC43 S	(897	CSKASS-		-RSATECLLEDKY	KUSDVGFVE	AYNNE
avianinfectiousbronchitisvirusS		SRRKRS-		Leedeleetsvi		
bovine coronavirus S		CNKVSS-		-RSATULLIFSKV		
 mouse hepatitis virus S 			PSAIR	GRSALIDLLFOR	KLSDVGEVE	I NNC
Consensus	(921	CAK SS		RSAIEDLLFDKV	KLSDVGFVE	AYNNC

					S	ection 2
L	(961)	961 .	970	980	990	100
human coronavirus OC43 S	(930)	GGAEI	RDITUVOS	KGIK FUPPL	LSENQISGY	LAAT
avianinfectiousbronchitisvirusS	(717)	APPLGFF	KULACARE	XNGLL LIBET	LTAEMQALKI	SSLV
bovine coronavirus S	(940)	GGAEE	RELECTOR	ANGLE AND PER	SENDISCL	LAAT.
mouse hepatitis virus S	(896)	改造の前※	RLLLGVOS	ENGIKA ELEV	SESOTSG	TGAT
Consensus	(961)	GGAE I	KDTICAÓS.	YNGIKVLPPL	LSENQISGYT	LAAT
					s	ection 2
	(1001)		,1010	.1020	,1030	104
human coronavirus OC43 S	(968)	ASLEPEW	THACVEL	YLNVAYRIIG	LAVIMOVESC	NOKE
avianinfectiousbronchitisvirusS	(757)	SMANGGI	TARGATER	HITTAKLIOTA	LUTTOSLILL	MOEK
bovine coronavirus S	(978)	ASLAPPW	SAAAGVEY	HILTALIÖTA PALLYCYALY	COUMDV 550	NEKT
mouse hepatitis virus S	(334)	HALL E. W.	DITTER GIVENIE	SUSVOXMENT	LAVA MINITES F	THE KIM
Consensus	(1001)	ASLFPPW	SAAAGVPF	YLNVQYRING	LGVTMDVLSC	NOKL
						ection 2
	(1041)	1041	1050	.1060	.1070	108
human coronavirus OC43 S avianinfectiousbronchitisvirus	(1008)	NA UNE	LYATER	DATHSZITVKT	TANVANARA	Y NEW T
avianinfectiousbronchitisvirusS	(797)	ASSIKA	ICHMUEUE	RETSLALOOF	DIVSKOSAT	TTET
bovine coronavirus S	(1018)	ANACONS	LGATOES	DATESLVKT	ANNARA	NNT.
bovine coronavirus S mouse hepatitis virus S	(974)	SISAP IN	LGAIODGE	DATINSALGET	SVNANAEZ	INNE
Consensus	(1041)	ANAFNNA	LGAIQEGF	DATNSALVKI	DAVVNANAE	LNNL
					S	ection 2
	(1081)	1081	.1090	.1100		
human coronavirus OC43 S	(1081) (1048)	1081 00% SNRF	.1090	,1100 E. L'SRLD' L'B		
human coronavirus OC43 S avianinfectiousbronchitisvirusS	(1081) (1048) (837)	1081 QQUSNRI ASUNKHE	1090 	,1100 E 1,5 R D D 1 D E E 1,7 O P D 2 TO		
human coronavirus OC43 S avianinfectiousbronchitisvirusS bovine coronavirus S	(1081) (1048) (837) (1058)	1081 GOLSNET ASUNKHE GOLSNET	1090 (1.1 #ASLT) (1.1 9VTC (3.1 SSL)	,1100 E USRUDTUB ETYQQFDITO FLEREDADE		
human coronavirus OC43 S avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S	(1048) (837) (1058) (1014)	OQUSNRE ASUNKNE DOUSNRE NOLSVRE	OF TREE	,1100 E. USRUDT DE STYCOP() FTC FILSRED TIE STETRIES VE	1110 E CIPETO N. 0000 1 1 1 O FIDE 12	112 GEGT GEGT GEGT
human coronavirus OC43 S avianinfectiousbronchitisvirus S bovine coronavirus S mouse hepatitis virus S Consensus	(1048) (837) (1058) (1014)	OQUSNRE ASUNKNE DOUSNRE NOLSVRE	OF TREE	1100 E PSRUDIDE ETYCOPIE ELSELUTE ELLSELDALE	1110 E CIPETO N. 0000 1 1 1 O FIDE 12	112 GEGT GEGT GEGT
human coronavirus OC43 S avianinfectiousbronchiltsvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(1048) (837) (1058) (1014)	OQUSNRE ASUNKNE DOUSNRE NOLSVRE	OF TREE	,1100 Ligskupide Ligopolijo Palskupis Tolkie Ellskudale	,1110 EAGLEETE N. OMBELET OFFERE EKEGLOGGE ANAQIORLIK	112 GART CRUS IGGLT IGRLT
	(1048) (837) (1058) (1014) (1081)	QCASNRI ASTINKINA EQIJANRI NOLSHAI QCLSNRF	AST GILSSVIC G. SSLE WSTU GAISASLQ	1140	1110 E ZIPETO NEOVORALET O FIRELO EKACIDES LO ANAQIDELIA S	112 CD ST CD SS IG S T IG R L T IG R L T iection 2
	(1048) (837) (1058) (1014) (1081)	QCASNRI ASTINKINA EQIJANRI NOLSHAI QCLSNRF	AST GILSSVIC G. SSLE WSTU GAISASLQ	1140	1110 E ZIPETO NEOVORALET O FIRELO EKACIDES LO ANAQIDELIA S	112 CD ST CD SS IG S T IG R L T IG R L T iection 2
human coronavirus OC43 S	(1048) (837) (1058) (1014) (1081) (1121) (1088)	QUESNA ASTINKINA DOLENA NOLENA QUESNA 1121	TASU, SL, SV, SSL, AST, GAISASLQ 1130 CLSDSTLV	1140 KDS/AQAMEK	1110 E T T T T T T T T T T T T T T T T T T T	112 FET CREST GEST GEST IGRLT JGRLT JECTION 2
human coronavirus OC43 S avianinfectiousbronchitisvirus S bovine coronavirus S	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098)	QUESNA ASTINKINA DOLENA NOLENA QUESNA 1121 TINAYUSO ESVLASA	T ASTO	1140 KESWAQEMEK SOORELETOR	1110 EVALUE EN TOUR EN	112 ICATT COUNTY ICATT I
human coronavirus OC43 S avianinfectiousbronchitisvirus S bovine coronavirus S	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098)	QUESNA ASTINKINA DOLENA NOLENA QUESNA 1121 TINAYUSO ESVLASA	T ASTO	1140 KESWAQEMEK SOORELETOR	1110 EVALUE EN TOUR EN	112 ICATT COUNTY ICATT I
human coronavirus OC43 S avlaninfectiousbronchitisvirusS bovine coronavirus S mouse hepatitits virus S	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098) (1054)	QUISNA ASTINKNE DOT SUR VOLSUR VOLSUR 1121 TUAXVSQ LIVEVIASA LIVEVIASA LIVEVIASA	TASTO	1140 KESKAQIMEK SOORELKIOK KESAAQAMEK KVSAAOSIEK	1110 ENTERNATION OF THE TOTAL O	112 105 T 105 T 105 T 105 T 116 Ection 2 116 EX T 117 T 117 T 117 T 117 T 117 T 117 T
human coronavirus OC43 S avlaninfectiousbronchitisvirusS bovine coronavirus S mouse hepatitits virus S	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098) (1054)	QUISNA ASTINKNE DOT SUR VOLSUR VOLSUR 1121 TUAXVSQ LIVEVIASA LIVEVIASA LIVEVIASA	TASTO	1140 KESWAQEMEK SOORELETOR	1110 NICONDESS NICON	112 GARTE GA
human coronavirus OC43 S avlaninfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098) (1054) (1121)	ASSINKINA ASSINKINA POLSINAT NOLSINAT OQLSINAT 1121 NAXYSO LINAYVSO LINAYVSO 1161	AS OF THE PROPERTY OF THE PROP	1140 KESWAGAMEK SORELHTOS KESWAGAMEK KVSBAGATEK KFSBAQAMEK	1110 LLLLE LE LE LE LE LE LE LE LE LE LE LE L	112 112 113 114 116 116 116 117 117 118 118 118 118 118 118
human coronavirus OC43 S avlaninfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098) (1054) (1121)	ASSINKINA ASSINKINA POLSINAT NOLSINAT OQLSINAT 1121 NAXYSO LINAYVSO LINAYVSO 1161	AS OF THE PROPERTY OF THE PROP	1140 KESWAGAMEK SORELHTOS KESWAGAMEK KVSBAGATEK KFSBAQAMEK	1110 LLLLE LE LE LE LE LE LE LE LE LE LE LE L	112 112 113 114 116 116 116 117 117 118 118 118 118 118 118
human coronavirus OC43 S avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(1048) (837) (1058) (1014) (1081) (1121) (1088) (877) (1098) (1054) (1121) (1161) (1128)	ASPINKNA ASPINKNA MOLSHAF NOLSHAF NOLSHAF LEVIA	GAISASLQ 1130 GISDSTIV KONSTIV 100 1100 GISDSTIV QLSDSTIV LIVENARY 510 1170 510 100 100 100 100 10	1140 KESAROTMEK SOORELETOS KESAROTMEK KVSAROTTES KFSAROTMEK 1180	1110 PROVINCE TO THE PROVINCE	112 112 123 123 123 124 136 146 147 147 147 147 147 147 147 147
human coronavirus OC43 S avlaninfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus human coronavirus OC43 S avlaninfectiousbronchitisvirusS bovine coronavirus S	(1048) (837) (1058) (1014) (1081) (1121) (1121) (1088) (877) (1098) (1054) (1121) (1161) (1128) (917) (1138)	ASTENDING ASTENDING BOLING MOLING HOLING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING	GALSASLQ AL	1140 KSBAAO MEK SQORELETOS KESA QAMEK KVSAAO TESA KFSAA QAMEK ,1180 GIYELETSYY GIVELETSYY	1110 TROUBLE TO THE TO	112 Part of the color of the co
human coronavirus OC43 S avianinfectiousbronchitisvirusS bovine coronavirus S mouse hepatitis virus S Consensus	(1048) (837) (1058) (1014) (1081) (1121) (1121) (1088) (877) (1098) (1054) (1121) (1161) (1128) (917) (1138)	ASTENDING ASTENDING BOLING MOLING HOLING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING LICENDING	GALSASLQ AL	1140 KSBAAO MEK SQORELETOS KESA QAMEK KVSAAO TESA KFSAA QAMEK ,1180 GIYELETSYY GIVELETSYY	1110 TROUBLE TO THE TO	ection 2 116 FTME FYST FYST FYST FYST FYST FYST FYST FYST

						Sec	ction 3
(1201)	1201	,1210		1220	,123		124
human coronavirus OC43 S (1168)	AGDRG		PKS	YEVNVNI	TWMYTGS	3YYY	E Pile
avianinfectiousbronchitisvirusS (957)	KPANAS	YAIVPA	NGR.	INTOVNO	SYYLWAR	MYM	RAL
bovine coronavirus S (1178)	AGDRG		PKSU	YEVNVNI	TWMF2GS	TYTYE	EPI
mouse hepatitis virus S (1134)			PKA'S	XEAODDO	ENKRIGS	SXXXP	EPT
Consensus (1201)	AGDRG	IA	PKSG	YFVNVNI	TWMFTGS	SYYYP	EPI
						Se	ction 3
(1241)	1241	.1250		,1260	,127	0	128
human coronavirus OC43 S (1202)	PINNVVVI	STOAVE	TKA	PYVMLN	CSIP-NEP	DEKEL	DO
avianinfectiousbronchitisvirusS (997)	AGDMAT	TSCOAN	VSV	NKTVIIT	FVDNDDE	DENDE	LSK
avianinfectiousbronchitisvirusS (997) bovine coronavirus S (1212)	ENVIEW	ATTE AUN	TKA	ROVMIN	STP-NEP	YKE	LDO
mouse hepatitis virus S (1168)	DENSTE	ISSCAVE	MEKA	PEVEUN	STP-NP	D K F	A NK
Consensus (1241)	ENNVVVI	MSSCAVN	YTKA	PDVMLN	CSIP NLP	DFKEE	LDO
	200000						ction 3
(1281)	1281	.1290		,1300	.131	0	132
human coronavirus OC43 S (1241)	FREEDING	THE STATE	D.Y	TNUPER	FTOVEMN	COPAT	K.V.T
avianinfectiousbronchitisvirusS (1037)	WNDTKH	LEDEDK	FN	YTYPI	IDSELDE	LOGVI	OGL
hovine coronavirus S (1251)	EKNOTS	PARTER	6	H.N. OTREAT	TODEMNI	LSHA	KVE
avianinfectiousbronchitisvirus (1037) bovine coronavirus S (1251) mouse hepatifis virus S (1207)	PENOTS	TAPPLE	DEEK	LNACLI	LTYEMNE	IODAI	KK
Consensus (1281)	FKNOTS	VAPDLSL	DY	INVTFL	DLO EMNR	IOEAI	KVL
					~ ~		ction 3
(1321)	1321	,1330		1340	135	0	136
human coronavirus OC43 S (1279)	DAYINE	KDEGTYE	YIVI	RENGMA	LECLAGY	AMUVI	THE
avianinfectiousbronchitisvirusS (1075)	DSLANDE	FKESILK	TAIR	CHRIST	ATAFATT	IFEL	LCW
bovine coronavirus S (1289)	OUTINE	KDIGTYE	YIVE	联系数据	LIGFAGV	AMT VI	EFF
bovine coronavirus S (1289) mouse hepatitis virus S (1247)	ESYDNE	KEVGTME	MIVE	TER SAY	LICUACY	AVCVI	FE
Consensus (1321)	OSYINL	KDIGTYE	YYVK	WPWYVW	LLIGLAGV	AMLVI	LFF
					1	Se	ction 3
(1361)	1361	1370		,1380	,139		140
human coronavirus OC43 S (1319 avianinfectiousbronchitisvirusS (1115	CCCTGC	ğ	街	SCFRE	GCCDDYT	GYCE	sví k
aylaninfectiousbronchitisvirusS (1115)	FFME	ČGCCCGC	FGIM	PLMSKO	GKKSSY T	TFDNI	DVVT
bovine coronavirus S (1329)	ecc to	ĕ −−−−		SOEKKE	GECDENT	GHQE.	LVIK
mouse hepatitis virus S (1287)	ccolec	Ë	S	CCFKE	SNOCDENG	GHQD	SIVI
	CCCTGC	G ·	T	SCFKKC	GGCCDDYT		
Consensus (1361)						90	ction 3
Consensus (1361							
Consensus (1361) (1401)		408			1 1		
) 1401 1	SEQ		0: 9918			
(1401 human coronavirus OC43 S (1350 ayianinfectiousbronchitisvirusS (1155) 1401 1) SHDD	SEQ SV SEQ	ID N	O: 9909	*		
(1401 human coronavirus QC43 S (1350) 1401 1) SHDD	SV SEQ	ID N	O: 9909 O: 9891	*	30	
(1401 human coronavirus OC43 S (1350 ayianinfectiousbronchitisvirusS (1155) 1401 1) \$HDD) QMRPKK) SHED	SV SEQ	ID N	O: 9909		30	

						- Section 15
	(589)	589	,600	.610	.620	630
human coronavirus OC43 S	(565)	NSOT	REQAPLENT	APSCTOCK	NICRUETTHO	Alexan Iraca
bovine coronavirus S	(575)	NPCTC	QPOAFLGVS	VDSCLOGORO	NICEPTERIO	/ Sommons
mouse hepatitis virus A59 S	(524)	KGĢIĢ	ANNSFIGUE	HUTCLVNER	OBSANTI ING	CHEST THE S
Consensus	(589)	N CTC	POAFLGWS	DSCLQGDRC	NIFANFILHD	INSGTTCS
						Section 16
	(631)		640	.650	.660	672
human coronavirus OC43 S	(607)	TOLOK	ANIDITLEV	WNYDIAGOL	ed Pitypyn As	Market A Balletin
bovine coronavirus S	(617)	PULLUK	VELTIDENS	ON NOTES FOR AT	COLTEVENAL	SEA TO SEA SEA
mouse hepatitis virus A59 S	(566)	TOIRL	PNTEVVTUT	eokydiisiac t	COSTEWNATED KAI	DESTRUCTION OF THE PARTY
Consensus	(631)	TDLQK.	ANTDIILGV	CVNYDLYGIT	GOGIFVEVNA	YYNSWON
						Section 17
	(673)	673	680	,690	,700	714
human coronavirus OC43 S	(649)	LUIDS	NOTE YOUR D	YTINRTEMER	SOYSIGNUSANI	ANSSER
bovine coronavirus S	(659)	LINDS	LOLLY CHARLE	YLTHRIFMIR	SCYSS PHYSIALS	LANSSET
mouse hepatitis virus A59 S	(608)	I THE DV	LEGIND FROM	LTINKIYTI		KDAPER
Consensus	(673)	LLYDS	NGNLYGFRD	YITNRTFMIR	SCYSGRVSAAL	HANSSEP
						Section 18
	(715)	715	720	730	740	756
human coronavirus OC43 S	(691)	AIR BY	LIKENSTER	SLUFOROPI	NETDEL LOTAL	ATCUYA
bovine coronavirus S	(701)	ALLERA	VCK-NI-5N	SITPOLOPI THSPOLOPI	aled Syletya	TADVSTS
mouse hepatitis vīrus A59 S	(650)	ABBYR	TENEST VINS	PULSTIEENPL	HYER STREET	ADNRED
Consensus	(715)	ALLFR	NIKCHYVFN	NSLSRQLQPI	NYFDSYLGCVV	NADNSTA
				-··· S1		∠Section 19
	(757)	757	,77	0 ← 78	30	798
human coronavirus OC43 S	(733)	ISVOI	TTTVGSGY.	VL SKNES	RGAITTCYRET	NEBPETY
bovine coronavirus S	(143)	SAAGA	からばかればみだ 式。	MADEUT KEES	RRSTURGYRDS	NEEDFOV
nouse hepatitis virus A59 S	(692)	EALPN	DERMCACL	VETSKSPRA	HRSVSTQ BLI	TELEXEP
Consensus	(757)	AVQT	CDLTVGSGY	CVDYSK RRS	RRSITTGYRFT	NEEPETV
						 Section 20
	(799)	799	810	820	.830	840
human coronavirus OC43 S	(775)	NSW5D	LEPVGC 11.10	OROLL SECTIO	830 CNMVEFFTES CNMBEFFTES	SKVQTES
bovine coronavirus S	(785)	NSWED	LEPAGGLI	I DIPSEEVE	Gnmber fors.	EVELED
mouse hepatitis virus A59 S	,	7.8.032	が作れて、海口は光明子は	台西西米的银行运动	THE PROPERTY OF THE PARTY OF TH	The state of the state of
Consensus	(799)	NSVNDS	LEPVGGLY	ZIQIPSEFTI	GNMEEFIQTSS	PKVTIDC
						- Section 21
	(841)	841	,850	860	,870	882
human coronavirus OC43 S	(817)	AATVE	DYAAGKSG	VEYGUECDS	ingititiyyei ingititiyyei	Tratto in
bovine coronavirus S	(827)	SATUE	DYBACKSO	VEIGEZUDE	INALLITEVNEL	LUTTOLO
mouse hepatitis virus A59 S	(116)	WALKELLO.	UNTARROOM	VETCHEUV	VNATANAMAN	SINMSTO
Consensus	(841)	AAFVC	DYAACKSOI	VEYGSECON	INAILTEVNEL	LDTTOLO

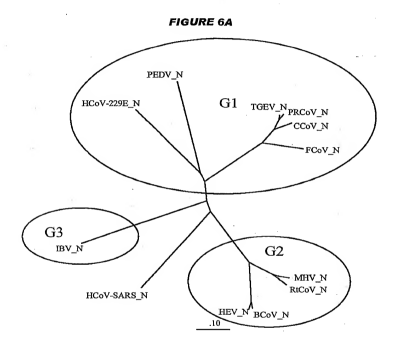


FIGURE 6B

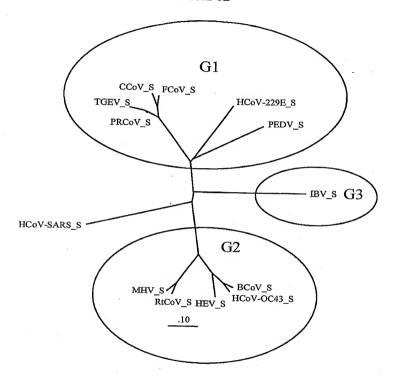


FIGURE 6C

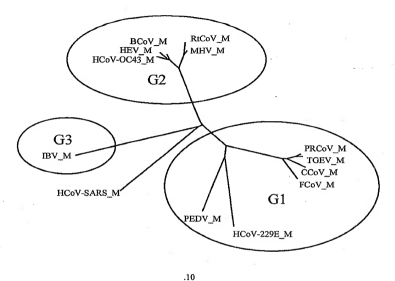


FIGURE 7

FIGURE 7A

SEQ	ID	NO:605	3MFVLLVAYALLH	12
SEQ	ID	NO:605	7MKKLFVVLVVMPLIYGDNFPCSKLTNRTIGNQWNLIETFLLNYSSRLPPNSDVVLGD	60
SEQ	ID	NO:606	1MRSLIYFWLLLPVLPTLSLPQDVTRCQSTTNFRRFFSKFNVQAPAVVVLGG	52
SEQ	ID	NO:606	5MFLILLISLPTAFAVIGDLKCTTVS-INDVDTGVPSIS	30
SEQ	ID	NO:606	9MLFVFILFLPSCLGYIGDFRCIQLVNSNGANVSAPSIS	10
SEQ	ID	NO:604	2MFIFLLFLTLTSGSDLDRCTTFDDVQAPNYTQHTSSMRG	20
SEO	ID	NO:607	2MLVTPLLLVTLLCALCSAVLYDSSS	27
_			i. ; ,	41
			., , ,	
SEO	ID	NO:605	3	
		NO:605		120
SEO	ID	NO:606	YLPSMNSSSWYCGTGIETASGVHGIFLSYIDSGQGFEIGISQEPFDPSGYQLYLH	10
		NO:606	5YY	TO.
		NO:606	YY	71
		NO: 604		1,1
		NO:607		19
226		210.007	IVIIIQSAFRFPSGWHLQG	46
			·	
SEO	TD	NO:605	3IAGCQTTNGLNTSYSVCNG	24
		NO:605	TTRNFNSAEGALICICKGSPPTTTTESSLTCNWGSECRLNHKFPICPSNSEANCGNMLYG	31
		NO:606	KATNGNTNAIARLRICQFPDNKTLGPTVNDVTTGRNCLFNKAIPAYMRDGKDIVVGITWD	180
		NO:606		10
		NO:606		123
		NO:604		131
		NO:607		T32
~-£		-10,000	ONINA AMIDDEL MANAGED DOCT AGITUPORA AMADETAMINA	88
			• •	
SEO	ID	NO:605		
		NO:605		240
SEQ	ID		LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYY	240
SEQ SEQ	ID ID	NO:605	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRRSCAMQYYYTPTYYMIN	214
SEQ SEQ SEQ	ID ID ID	NO:605 NO:606 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPYYYMLN 5STFVNTSYSVVVQPHTTILGNKLQGFLEISVCOYIMCBYDNT	214 171
SEQ SEQ SEQ SEQ	ID ID ID ID	NO: 605 NO: 606 NO: 606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPTYYMLN	214 171 165
SEQ SEQ SEQ SEQ SEQ	ID ID ID ID	NO: 605 NO: 606 NO: 606 NO: 604	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRRSCAMQYVYTPTYYMLN STFVNTSYSVVVQPHTTILGMKLQGFLEISVCQYTMCBYPNT	214 171 165
SEQ SEQ SEQ SEQ SEQ	ID ID ID ID	NO: 605 NO: 606 NO: 606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRRSCAMQYVYTPTYYMLN	214 171 165
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SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID	NO:605 NO:606 NO:606 NO:606 NO:604 NO:605	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPYYYMLNSTFVNTSYSVVVQPHTTLIGKKLQGFLSISVCQYTMCEYPNT	214 171 165 158
SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID	NO:605 NO:606 NO:606 NO:606 NO:604 NO:607	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRRSCAMQVVYPPTYYMLN	214 171 165 158 74
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID	NO: 605 NO: 606 NO: 606 NO: 606 NO: 607 NO: 605 NO: 605	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVTPTYYMLN	214 171 165 158 74 300 274
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID ID ID ID ID ID I	NO:605 NO:606 NO:606 NO:606 NO:604 NO:605 NO:605 NO:605 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDUSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPTYYHLNSFVNTSSYSVVVQPHTTLIGKKLQGFLBISVCQYTMCEYPNT	214 171 165 158 74 300 274 230
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID	NO: 605 NO: 606 NO: 606 NO: 606 NO: 607 NO: 605 NO: 605 NO: 606 NO: 606 NO: 606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYPPTYYMLN STFVNTSYSVVVQPHTTLIGNKLQGPLEISVCQYTMCEYPNTSLFGYTSYTVVIEPYN	214 171 165 158 74 300 274 230
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID	NO:605 NO:606 NO:606 NO:606 NO:607 NO:605 NO:605 NO:606 NO:606 NO:606 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYPPTYYMLN	214 171 165 158 74 300 274 230 225 217
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID	NO: 605 NO: 606 NO: 606 NO: 606 NO: 607 NO: 605 NO: 605 NO: 606 NO: 606 NO: 606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVTPTYYMLN	214 171 165 158 74 300 274 230 225 217
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID	NO:605 NO:606 NO:606 NO:606 NO:607 NO:605 NO:605 NO:606 NO:606 NO:606 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYPPTYYMLN	214 171 165 158 74 300 274 230 225 217
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID	NO:605 NO:606 NO:606 NO:606 NO:607 NO:605 NO:605 NO:606 NO:606 NO:606 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPTYYHLNSTPVNTSYSVVVQPHTILIGKKLQGFLEISVCQYTMCEYPNT	214 171 165 158 74 300 274 230 225 217 131
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID	NO:605 NO:606 NO:606 NO:606 NO:605 NO:605 NO:605 NO:605 NO:606 NO:606 NO:606 NO:606 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPTYYMLN SFWYNTSYSVVVQPHTTLIGKKLGGPLEISVCQYTMCEYPNT	214 171 165 158 74 300 274 230 225 217 131
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID ID	NO:605 NO:606 NO:606 NO:604 NO:605 NO:605 NO:606 NO:606 NO:606 NO:606 NO:606 NO:606 NO:606	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNPVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYPPTYYMLN	214 171 165 158 74 300 274 230 225 217 131
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID ID ID ID ID ID I	NO:605 NO:606 NO:606 NO:606 NO:605 NO:605 NO:605 NO:606 NO:606 NO:607 NO:607	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPYYYMLN STFVNTSYSVVVQPHTILIGKKLQGFLEISVCQYTMCEYPNT	214 171 165 158 74 300 274 230 225 217 131 360 332
SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ	ID ID ID ID ID ID ID ID ID ID ID ID ID I	NO:605 NO:606 NO:606 NO:606 NO:606 NO:605 NO:605 NO:605 NO:606 NO:606 NO:606 NO:606 NO:605 NO:605 NO:605	LQWFADEVVAYLHGASYRISFENQWSGTVTFGDMRATTLEVAGTLVDLWWFNFVYDVSYYNDRVTVFADKIYHFYLKNDWSRVATRCYNRSCAMQVVYTPTYYMLN STFVNTSSYSVVVQPHTTLIGKKLGGPLEISVCQYTMCEYPNT	214 171 165 158 74 300 274 230 225 217 131 360 332 286

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SEO	ID	NO:6057	VFSLNTTGGVTLEISCYTVSDS-SFFSYGEIPFGVTDGPRYCYVHYNGTALKYLGTL	
		NO:6061	IVLHTALGTNLSFVCSNSSDPHLAIFAIPLGATEVPYYCFLKVDTYNSTVYKFLAVL	
		NO:6065	CKSDFMSEIKCKTLSIAPSTGVYELNGYTVOPIADVYRRIPNLPDCN-IEAWLNDKSVPS	
		NO:6069	CASSYTSEIKCKTQSMLPSTGVYELSGYTVQPVGVVYRRVANLPACN-IEEWLTARSVPS	
		NO:6042	CSQNPLAELKCSVKSFEIDKGIYQTSNFRVVPSGDVVR-FPNITNLCPFGEVFNATKFPS	
		NO:6072	FKAGGPITYKVMREVKALAYFVNGTAODVILCDGSPRGLLACOYNTGNFSDGFYPFTNSS	
PPQ	ID	10.0072		231
C PO	TD	NO:6053	PKTVREFVISRTGHFYINGYRYFTLGNVEAVNFNVTTAETTDFCTVALASYADVLV	242
		NO:6057	PPSVKEIAISKWGHFYINGYNFFSTFPIDCISFNLTTGDSDVFWTIAYTSYTEALV	
		NO:6061	PPTVREIVITKYGDVYVNGFGYLHIGLIDAVTINFTGHGTDDDVSGFWTIASTNFVDALI	
		NO:6065	PLNWERKTFSNCNFNMSSLMSFIQAYSFTCNNIDAAKIYGMCFSSITIDKFAIPNG	
		NO:6069	PLNWERKTFQNCNFNLSSLLRYVQAESLFCNNIDASKVYGRCFGSISVDKFAVPRS	
		NO:6042	VYAWERKKISNCVADYSVLYNSTFFSTFKCYGVSATKLNDLCFSNVYADSFVVKGD	
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SEQ	ID	NO:6061	EVQGTSIQRILYCDDPVSQLKCSQVAFDLDDGFYPISSRNLLSHEQPISFVTLPSFNDHS	509
SEQ	ID	NO:6065	RKVDLQLGNLGYLQSFNYRIDTTATSCQLYYNLPAANVSVSRFNPSTWNRRFGFTEQS	459
SEQ	ID	NO:6069	RQVDLQLGNSGFLQTANYKIDTAATSCQLHYTLPKNNVTINNHNPSSWNRRYGFNDAG	
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SEO	ID	NO:6053	FIVLYVDFKPQ-SGGGKCFNCYPAGVNITLANFNETKGPLCVDTSHFTTKYVAVYAN	356
SEO	ID	NO:6057	IVNITIGLGMKRSGYGQPIASTLSNITLPMQDHNTDVYCIRSDQFS-VYVHSTCK	
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		NO:6072	VFKGRAT	
CEO	TD	NO:6053	VGRWSASINTGNCPFSFGKVNNFVKFGSVCFSLKDIPGGCAMPIVA	402
		NO:6057	SALWDNIFKRNCTDVLDATAVIKTGTCPFSFDKLNNYLTFNKFCLSLSPVGANCKFDVAA	
		NO:6061	SYGYVSKSQDSNCPFTLQSVNDYLSFSKFCVSTSLLAGACTIDLFG	
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		NO:6053	NWAYSKYYTIGSLYVSWSDGDGITGVPQPVEGVSSFMNVTLDKC	446
		NO:6057	-RTRTNEQVVRSLYVIYEEGDNIVGVPSDNSGVHDLSVLHLDSC	
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		NO:6069	KGCICANNSFIGWSHDTCLVNDRCQIFANILLNGINSGTTCSTDLQLPNTEVVTGIC	
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SEQ	ID	NO:6053	TKYNIYDVSGVGVIRVSNDTFLNGITYTSTSGNLLGFKDVTKGTIYSITPC	497
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SEQ	ID	NO:6065	VNYDLYGITGQGIFVEVNATYYNSWQNLLYDSNGNLYGFRDYLTNRTFMIRSC	685
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		NO:6042	VNFNFNGLTGTGVLTPSSKRFQPFQQFGRDVSDFTDSVRDPKTSEILDISPC	033
		NO:6072	VDYNIYGRTGQGFITNVTDSAVSYNYLADAGLAILDTSGSIDIFVVQGEYGLNYYKVNPC	2/0
DDQ		110.0072		493
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		NO: 6057	DVSAQAAVIDGTIVGAITSINSELLGLTHWTTTPNFYYYSIYNYTNDRTRGTAIDSND	542
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		NO:6065	YSGRVSAAFHAN-SSEPALLFRNIKCNYVFNNTLSRQLQPINYFDSYLGCVVNADNSTS	741
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PPÕ	ıD	NO.0072	VANCPYVSYGKFCIKPDGSIATIVPKQLEQFVAPLFNVTENVLI	588
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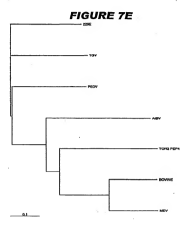
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SEQ ID NO:6061 SEQ ID NO:6065 SEQ ID NO:6069 SEQ ID NO:6042 SEQ ID NO:6072	CGCIGCLGSCCHSICSRRQFENYEPIEKVHVH 1450 CGCCGCCGACFSGCCRGPRLQPYEAFEKVHVQ 1384 G-TSCFKKCGGCCDYTGHQELVIKTSHED- 1364 G-SCCFKKCGNCCDEYGGHQDSIVIHNISSHED- 1326 CSCLKGACSCG-SCCKFDEDDSEPVLKGVKLHYT- 1255 CGCCCGCFGIMPLMSKCGKKSSYYTTFDNDVVTEQYRPKKSV 1164	
	FIGURE 7B	
SEQ ID NO:6054 SEQ ID NO:6058 SEQ ID NO:6045 SEQ ID NO:6045 SEQ ID NO:6066 SEQ ID NO:6066 SEQ ID NO:6066 SEQ ID NO:6068 SEQ ID NO:6068 SEQ ID NO:6068 SEQ ID NO:6068 SEQ ID NO:6068 SEQ ID NO:6068 SEQ ID NO:6066		54 59 52 57
	FIGURE 7C	
SEQ ID NO:6055 SEQ ID NO:6063 SEQ ID NO:6059 SEQ ID NO:6067 SEQ ID NO:6070 SEQ ID NO:6074		24 60 27 33 21 28
SEQ ID NO:6055 SEQ ID NO:6063 SEQ ID NO:6059 SEQ ID NO:6067 SEQ ID NO:6070 SEQ ID NO:6046	GWANTLITIETVILOPGHYKYSRLFYGLKMLVLWLLWPLVLALSIPDTWANWDSN-WAFVA TWNILITILLVUQYGHYKYSVFLYGVKWAILWILWPLVLALSIPDAWASFQN-WVFFA SWSILLIVEITVLQYGRPQFSWFYYGIKMLIMWLLWPVVLALTIFNAYSEYQVSRYVMFG SLGIILLFITVILOPGYTSRSMFYYVLKWYLIWLMWPLTITLTIFN-CYYALN-NVYLG SLGIILLFITTILQFGYTSRSMFIYVVKMIILWLMWPLTIVLCIFN-CVYALN-NVYLG VIGFLFLANIMLLQFAYSNRNRFLYIIKLVFLWLLWPVTLACFVLA-AVYRIN-WVTGG	83 120 84 90

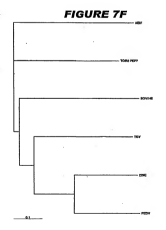
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	ID NO:6063	FSILMACITLMLWIMYFVNSIRLWRRTHSWWSFNPETDALLTTSVM-GRQVCIPVLGAPT	1/12
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SEQ	ID NO:6074	AAIILTVFACLSFVGYWIQSIRLFKRCRSWWSFNPESNAVGSILLTNGQQCNFAIESVPM	145
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		FIGURE 7D	
		FIGURE 7D	
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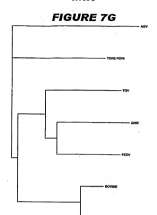
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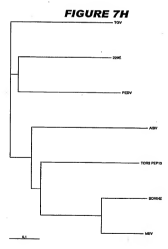
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		NO:6064		49
		NO:6060		15
SEQ	ID	NO:6068	8AGSRSRANSGNRTPTSGVTPDMADQIASLVLAKLGKDAAKP 2	51
SEQ	ID	NO:6071	1PASTVKPDMAEEIAALVLAKLGKDAGQP 2	52
SEO	ID	NO:6051	1nstpgssrgnsparmasgggetalalllldrlnqleskv 2	22
		NO:6075	5APSREGSRGRR 2	.33
U		1.0.0073	ALONDONGKY SDSGDDPTAKAWIIĞDÖ S	.12
ano	70	*** ***		
		NO:6056		65
		NO:6064		00
		NO:6060	ORSKSKERSNSKTRDTTPKNENKHTWKRTAGKGDVTRFY 2	53
SEQ	ID	NO:6068	BQQVTKQTAKEIRQKILNKPRQKRSPNKQCTVQQCF 2	86
SEQ	ID	NO:6071	KQVTKQSAKEVRQKILN	87
SEO	ID	NO:6051	L SGKGQQQQGQTVTKKSAAEASKKPRQKRTATKQYNVTQAF 2	75
		NO: 6075		./3
520		110.00,5		47
			:* *:	
CEO	TD	NO:6056	CDDDIDY MOCACINA	
				14
		NO:6064		49
		NO:6060		02
		NO:6068		43
SEQ	ID	NO:6071	GKRGPNQNFGGSEMLKLGTSDPQFPILAELAPTVGAFFFGSKLELVKKNSGGADE 3	42
SEQ	ID	NO:6051	GRRGPEQTQGNFGDQDLIRQGTDYKHWPQIAQFAPSASAFFGMSRIGMEVTP 3	27
SEO	ID	NO:6075	GPRTKGK-EGNFGDDKMNEEGIKDGRVTAMLNLVPSSHACLFGSRVTPKLQL2	00
			* * *** * : * . : : .	90
CEO	TD	NO:6056	MINITE III III III III III III III III III	
		NO:6064		49
				01
		NO:6060		43
		NO:6068		97
SEQ	ID	NO:6071	PTKDVYELQYSGAVRFDSTLPGFETIMKVLNENLNAYQKDGGADVVSPKPQRKGRR 3	98
SEQ	ID	NO:6051	SGTWLTYHGAIKLDDKDPQFKDNVILLNKHIDAYKTFPPTE3	68
SEO	ID	NO:6075	DGLHLRFEFTTVVPCDDPQFDNYVKICDQCVDGVGTRPKDDEPKPKSRSSSRPATRG 3	
_				22
			the state of the s	
SEO	TD	NO:6056	OHDLINDCALEDADCOMCD AMARDINDRIGHTED TIDELS	
		NO:6064		9
				6
		NO:6060		5
		NO:6068		8
SEQ	ΙD	NO:6071	QAQEKKDEVDNVSVAKPKSSVQRNVSRELTPEDRSLLAQILDDGVVPDGLEDDSNV 45	4

SEQ ID NO:6051	-PKKDKKKKTDEAQPLPQRQKKQPTVTLLPAA	399
SEO ID NO:6075	NSPAPROORPKKEKKLKKODDEADKALTSDEERNNAOLEFYDEPKVINWGDAALGENEL	414









		Section 1
(1) 1 10 20 artial 5'UTR 161- (1) TATTAAAATCTTATTGTTGCTGG:	.30	4.
artial 5'UTR 161- (1) TATTAAAATCTTATTGTTGCTGG	TATCACTGCTT	STTTTGC
cov-OC43 5'UTR (1)		
Consensus (1)		
		— Section 2
(43) 43 50 60 artial 5'UTR 161- (43) GTGTCTCACTTTATACATCTGTT(70	<u> </u>
aniai 5UIR 161- (43) GTGTCTCACTTTATACATCTGTT	GCTTGGGCTAC	CTAGTGT
coV-OC43 5'UTR (1)		
Consensus (43)		
		C
(85) 85 90 100 artial 5'UTR 161- (85) CAGCGTCCTACGGGCGTCGTGGC		Section 3
05) 05 90 100	,110	12
cov-oc43 5'UTR (1)	TGGTTCGAGTG	CGAGGAA
povine CV 5'UTR (1)		
O		
(127) 127 140 artial SUTR 161- (127) CTCTGGTTCMCCTAECGGTEGGC		Section 4
(127) 127 140	150	16:
artial 5'UTR 161- (127) CTCTGGTTCMTCTACCGCTAGGCC	Comon Carcas	n Chin Chin
CoV-OC43 5'UTR (1)	CCEPECTENCON.	- TO COCO
bovine CV 5'UTR (1)GATTCCGAGGGTTTT	George Caroca	Tecccc
Consensus (127) GATTG GAGCGATTT	GCGTGCGTGCA	TCCCGG
	· · · · · · · · · · · · · · · · · · ·	- Section 5
artial SUTR 161-(127) CTCTGGTTCMCTAGCGGTTGGC CO42 SUTR (1)	200	21
artial 5'UTR 161- (169) TECAGACGTACCGCTTCTGTTGT	GTCAAAEAC	SGGGTCAC
CoV-OC43 5'UTR (33) INTCA OTGATICE COTO	TAGATCTTTTT	STARTOTA
bovine CV 5'UTR (33) TTCACTCATCTCTTGT:	TAGATOTTTO	ATAATCT
Consensus (169) TTCA CTGATCTCTTGT	TAGATCTTTTC	STAATCT
		Section 6
(211) 211 220 230 artial S'UTR 161- (209) CTGCCCCCACATACCT CTAAGGG COV-OC43 S'UTR (68) AAGTTTATAAANACATCCACCCC	240	25
artial 5'UTR 161- (209) CTCCCCCCCCCTTACCTCTAGGG	CTTTTGAGCCT	AGCGTTG
CoV-OC43 5'UTR (68) AACTTTATAAAAAACATCGACTCG	CTGTAATCTAT	SCTEGEG
DOVING CV 5 UTR (68) AACETTAAAAAAAAAACAECACTCC	CTGTAGTCTATO	SCCTGTG
Consensus (211) AACTTTATAAAAACATCCACTCC	CTGTAGTCTAT	SCCTGTG
(253) 253 260 270 artial 5'UTR 161- (251) SCTACSTICTICGCATARGET CGG		Section 7
(253) 253 260 270	280	29
aniai 5UTR 161- (251) GCTACGTTCTCGCATAAGGTCGG	CTATACGACGT	TTGTAGG
COV-OC43 5'UTR (110) GCGTAGATTTTTCATAGTGGTGT bovine CV 5'UTR (110) GCGTAGATTTTTCATAGTGGTGT	TTATATT-CAT	TTCT-GC
Consensus (253) GCGTAGATTTTTCATAGTGGTGT	CHADAPI -CAT	THCT-GC
Conscissed (200) GCGTAGATTTTTCATAGTGGTGT		
		Section
(295) 295 300 310	320	33
partial 5'UTR 161- (293) GGTAGTGCCAAACA'ACCCCTGAG	GTGACAGGITC	TGCTGGT
(295) 295 300 310 partial SUTR 181- (293) GGTAGTGCBAAACAAGCCTTGAGCCV-OC43 SUTR (150) GTTAGCAGCTTTCAGCCAGGGC	GTGTTGTATCC	TAGGG
povine of a contract the care of the care	GIGIIGIATUU	TAGGGG
Consensus (295) GTTAACAGCTTTCAGCCAGGGAC		
	200	
(337) <u>337 </u>	300	373
Q ID NO: 9910 - (333) TTTAGTGAGCAGACATACAATAG Q ID NO: 9919 - (189)AGTG-GCCCCATAGGTC	ACAGI GACAAC	ATG
Q ID NO: 9892 (189)AGTG-GCCCACCCATAGGTC	ACAATG	
Consensus (337) AGTG GCCCACCCATAGGTC	DOUNTRILLE	-

		SEQ ID NO:
F1: $AT \frac{CTT}{TGC} G \frac{C}{A} G \frac{GT}{CG} A \frac{GGC}{TTT} G \frac{G}{C} GTG$	(136-154 nt)	6021
F2: $GTG_C^TGTG_C^GAT_C^{AG}CC_G^ACTTCA$	(152-172 nt)	6022 -
F3: CTTCAC $\frac{G}{T}G\frac{T}{A}TCT\frac{G}{C}TTGT\frac{GT}{TA}GA$	(168-195nt)	6023
R1: $AG_G^A \frac{ACCTGT}{TACAA}CAC \frac{C}{G}TC \frac{AGG}{CCT}GG \frac{T}{C}TG$	(307-329nt)	6024
R2: $AAA_T^C G_{AA}^{CG}TATA_{AA}^{GC}C_{AC}^{GA}C \frac{CT}{AC}TATG$	(265-288nt)	6025
R3: CGAC AT TATG CGA GA AC GTA GCCCA	(250-274nt)	6026

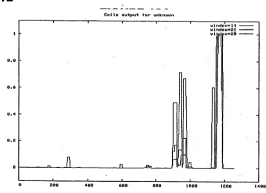
								Section	
	(1)	1		,10		20		5	36
avian IBV 3'UTR (NC_001451) 27103-	(1)	GTA	CATA	ATG	ACCT	STTGTTT	CCTGG	TACATTT	T
HCoV-OC43 3'UTR partial	(1)								-
bovine CV 3'UTR									-
Consensus	(1)							- " -	
**************************************								Section	
	(37)	37			50		60	7	72
avian IBV 3'UTR (NC_001451) 27103-	(37)	GTT	AAACA	CTAT	TTCTC	STGCTTT	CCTAT	CAATTAT	
HCoV-OC43 3'UTR partial bovine CV 3'UTR	(1)								_
Consensus									-
Consensus	(37)							Section	•
	/72\	73	. 0			00		Gecuon	20
avian IBV 3'UTR (NC_001451) 27103-	(73)	ACAC	CCAT	TCAT	PTCTCI	TOTATOT	TONNE	ACTTA AC	50
HCoV-OC43 3'UTR partial	(1)								_
bovine CV 3'UTR	(1)								_
Consensus	(73)								
								Section	
avian IBV 3'UTR (NC_001451) 27103-	(109)	109			20	,130)	14	44
avian IBV 3'UTR (NC_001451) 27103-	(109)	TTC	TTTT	GTT	CTTT'	TGCTTA	TTGTA	TTGTTGC	Т
HCoV-QC43 3'UTR partial	(1)								_
bovine CV 3'UTR									-
Consensus	(109)							Section	_
avian IBV 3'UTR (NC_001451) 27103-	(145)	145	150		,16	0	170	18	30
HCoV-OC43 3'UTR partial	(145)	GTG	STTTY	TAT	FGTTG!	FGATTCT	CATTE	ETTEG	問
bovine CV 3'UTR	(1)						TAAGA	GAALGAA	7
Consensus								GAATGAA	
	(170)							Section	
	(181)	181		.190		200			16
avian IBV 3'UTR (NC_001451) 27103-	(179)	TTTT	TCG			ATAGE	AGAGT		
HCoV-OC43 3'UTR partial	(14)	CIT	T-G	CGG	CACCTO	GTGGT	ACCCC	TC-GCAC	ď
bovine CV 3'UTR	(11)	CTI	4T-G	CGG	CACCTO	GTGGTA	ACCCC	TC-GCAC	G
Consensus	(181)	'CTT	AT GT	CGG	CACCTO	GGTGGT	ACCCC	TC GCAG	G
								Section	
avian IBV 3'UTR (NC_001451) 27103-	(217)	217			230	210- 311	240	2:	52
avian IBV 3'UTR (NC_001451) 27103-	(215)	ATA	SGCAT	GTA	SCTTG/	ATTACCI	ACATG	TCTATEC	c
HCoV-OC43 3'UTR partial bovine CV 3'UTR	(48)	AAA	GTCGC	G		AT,AAGGC	AC-TC	TCTATCA	LG,
bovine CV 3'UTR	(45)	AAA	GTCGC	- G		ATAAGGC	AC-TC	TCTATCA	řG

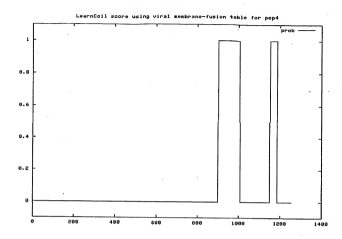
FIGURE 10 (contd.)

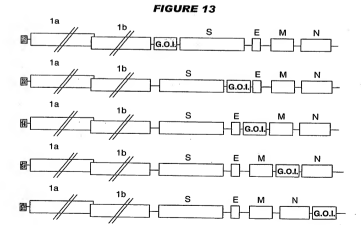
(05)	N 252	200			8	Section 8
avian IBV 3'UTR (NC_001451) 27103- (251) 253	260	35:2	70	W	288
HCoV-OC43 3'UTR partial (76) Andrea	HAARGIC	TAATCT	GTCTAC	TTAGTAG	CCTGG
bovine CV 3'UTR (73) anticc	- AGTC	TIGCIC	CIATAA	raga rag	AG
Consensus (253	AATGG	TCTC	TTTGCTG	CTATAA	BAGATAG	
	, mil 00.	1610	.116016	CTATAA		
/280	289	2	300	310		Section 9
avian IBV 3'UTR (NC_001451) 27103- (287	AMACC	A STICCHERON	Chicle cin	310		324
HCoV-OC43 3'UTR partial (107	AAGGT	PATECA	1 C 7 C T 7 T	TAGATT	LIANTER	AGTIT
bovine CV 3'UTR (104	AAGGT	PATAGGA	CACTAT			AGUTG AGUTG
Consensus (289	AAGGT	PATAGCA	GACTAT	ACATT		AGTTG
				MONII		ction 10
(325	325	330	340		250	
avian IBV 3'UTR (NC_001451) 27103- (323) AATTE	PAGTT	AGTTTA	AGTTAG	A CONTRACTOR	COLUMN TO THE REAL PROPERTY.
Consensus (325	AAAGT	TTGTGT	GGTAAT	GTATAGT	GTTGGA	GAAAG
					Se	ction 11
(361	361	370		380		396
avian IBV 3'UTR (NC_001451) 27103- (358	TATAA	GATGCC	AGTGCC	GGGCC	C-GCG	AGTAC
11004-0043 3 0 1K partial (1/5	I THE - IA A 2	1000 T		CONTRACTOR NO	im makes or or	21000
50VIII6 0V 301K (172		GACT	TGCG	GAACTA	TTGCCG	ACAAC
Consensus (361	TG AA	AGACT	TGCG	GAAGTAA	TTGCCG	ACAAG
					Se	ction 12
avian IBV 3" ITP (NC 001451) 27100 (397	397	1. / 1/6	410	,420	0	432
avian IBV 3'UTR (NC_001451) 27103- (393	GATOG	GGGTAC	AGCACT	AGGACGC	CCATTA	SGGGA
HCoV-OC43 3'UTR partial (206	TGCCC	AAGGGA	AGAGCC	ACCACG-		YSTTA
bovine CV 3'UTR (203 Consensus (397	TCCCC	AGGGGA	AGAGEC	ACCATG-		AGTTA
	IGCCCA	MGGGGA.	AGAGCC	AGCACG		AGTTA
1433	433	440	45		Se	ction 13
avian IBV 3'UTR (NC_001451) 27103- (429)	ACMCEN	MAN THE	70 7 7 7 7 7 T	mm's a citizen	GO CONTRACTOR E	468
HCoV-OC43 3'UTR partial (238)	CCACCC	ACTAAN	MACMAA	THAAGIII	AAGTTT	AIT-T
DOVING CV 3 O 1 K (235)	GUATEE	ACTAAT	TAGTAA	A TIC A A MC	7 A. C. m m. s	Comment and
Consensus (433	CCA CC	AGTAAT	TAGTAA	ATCANTO	AAGTTA	* + 11A11
				HIGHAIG	Se	tion 14
(469)	469	48	30	490		
avian IBV 3'UTR (NC_001451) 27103- (462)	GGCTAA	GTATAG	TTAAAA	mmm n m n o	CCTACT	
11C0V-0C43 301R partial (2/4)	GGCCAA	TRECAM	CAATCA	3		IIAGA
bovine CV 3'UTR (271)	GGCCAA	TTGGAA	GAATCA	E		
2				-		
					Sec	tion 15
(505)	505	513			560	
avian IBV 3'UTR (NC_001451) 27103- (498)	GTTAGA	GCD .		NO: 9911		
HCoV-OC43 3'UTR partial (293)		:		NO: 9920		
bovine CV 3'UTR (290)			CEO ID 1	NO: 9893		
Consensus (505)		'	DEG ID I	NO: 9893		

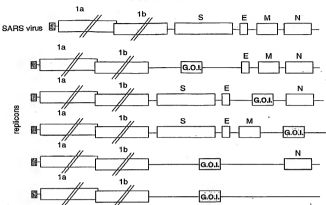
FIGURE 11

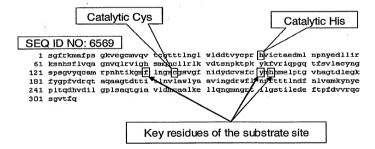
		SEQ ID NO
F-1 TCTATC GCC AGA GGATGTCT	(245 ~ 265 nt)	6027
F-2 TTAGTT $\frac{1}{G}$ AA $\frac{TT}{AG}$ TTT $\frac{A}{T}$ GT $\frac{T}{G}$ T $\frac{A}{G}$ GT	(318 ~ 339 nt)	6028
F-3 TAGTGTT $\frac{A}{G}$ GAG $\frac{T}{A}$ A $\frac{G}{A}$ GT $\frac{A}{G}$ TAAAGA	(346 ~ 368 nt)	6029
R-1 A $\frac{A}{C}$ TT $\frac{G}{A}$ GCCATA $\frac{A}{T}$ T $\frac{T}{A}$ AACTT	(458 ~ 476 nt)	6030
$R-2 \ ACTAA \frac{TTAC}{AATT} T \frac{G}{A} G \frac{C}{T} \frac{GG}{CT} T \frac{AA}{CC} C \frac{T}{C} TAA$	(426 ~ 448 nt)	6031
		6032
$H-3T \frac{TG}{AC}TC \frac{G}{C}GC \frac{AA}{G}T \frac{TA}{GG}C \frac{TT}{CC} \frac{C}{G}GCA$	(375 ~ 395 nt)	6033











							Section 1
100	(1)	1	,10	20	.3	04	0 51
avian IBV nsp2	(1)	SCEKKEVS	PESAVEK	CIVSVEY	RGNNLHGL	WLGDELYCPF	HVEGKFEG
MHV nsp2	(1)	SCHUKMVS	EMSK?EP6	TVSVTV	CHMPT. HOLL	WINDRWSCOR	WYTO COMPUMD
SARS nsp2	(1)	SCFRKMAF	PEGK VEG4	CHECKER	GTTTLNGL.	พร.ทฤหันของร	UT CHARRET AT
BCoV nsp2	(1)	SCHUKNAN	P#SKVEP(CIVSVTY	GNMTLNGL	WEDDKYYCPE	VICSASOMEN
Consensus	(1)	SGIVKMVS.	PSSKVEP	CIVSVTY	GNMTLNGL	WLDDTVYCPE	HVICSAADMIN
		····					Section 2
	(52)		30	.70	,80	90	102
avian IBV nsp2	(49)	DOMNORLN	LANNHEFI	TTOOHG	VEL NVESR	REFGAULELO	TAMANAETPKY
MHV nsp2	(52)	PDYPMLLC:	RVTSSDF	WHEGR -	WOLLD'MSY	OMORCOLVIA	MATERIAL DATE TO LANCE
SARS nsp2	(52)	PNYEDLLI	RKSWHSFI	VOAGN-	VOLR'	SMONCHERLK	VDTSNERTERY
BCoV nsp2	(52)	PDYTHLLC	RVTSSDFT	WAFDR -	MST.TI'MSY	OMOGCALVILT	VTLONSTPKY
Consensus	(52)	PDY NLLC	RVTSSDF	VLSGR	VSLTVMSV	OMOGCILLVI	VTLQNPKTPKY
					**************************************	Sugocupier	Section 3
30	(103)	103 .11	0	120	130	140	152
avian IBV nsp2	(100)	KEIKANCG	DEFTEACH	VGGRVV	GLEEPUPME	SMCOTERS	AGECSSYCHNE
MHV nsp2	(102)	SEGVVERG	ETFTYLA	YNGRPO	GARHUTER	SHITTEGER	CCSCCSVC
SARS nsp2	(102)	KEVRROPG	OF FEVER	VNCSTES	COMOCAME	purior record	NGSCUSVGENZ
BCoV nsp2	(102)	TEGVVKPG	EMPTUT.AT	VICKO	CVARALL	CENTRACET	CCSCCSVGY
Consensus	(103)	KEGVVKPG	FOR POTIL DE	VNGSDO	CAPHUTAD	CORMINCORT ASSETTACOS	CGSCGSVGFVI
- 3/4	1,			1111001.2	OAPH I INK	SOUTTVOOLD	Section 4
	(154)	154 .160		.170	.180	.190	
avian IBV nsp2	(151)	EKGUVNER	VERSIER L. E. L.	Militario	PORMORE	COMPERMIN	RVPPONLVTNN
MHV nsp2	(153)	TERSUMEY	de la Ser Er S	and Aug 2	Threeway	COUNTRACTOR	LEACDALOLAH
SARS risp2	(153)	Dynevsec	MASSEL NET.	oriettes a co	approus .	SELUDIAD AM	AAGTOTTITEN
BCoV nsp2	11531	MCDCURRY	101 101 101 10	MCCAMO	anguerr.	GTWK DKG LYG	TEAODAIGEAN WYGATHATAER
Consensus	11541	DCDCAKEA	MUOTET	mccume	DEFICATE I	GE TYPH CARG	LPVQDYTQTVN
	,,,,		MINGELLE	LOCALG	IDU GDEI	GET ADMOAAA	Section 5
	(205)	205 210		20	230	240	255
avian IBV nso2	(202)	MUAMILYAN			Trio mentr car	POSSERVE COM	GEMPFSMST
MHV nsp2	(204)	WWANTERS	CEMBCM-	TO DE LOS	NO CHECCE	Continue william	GESSEKADL
SARS nso2	(204)	WHAMILY C.	TMCDP-		DE EUCENE	CEL II O IN STREET	HEPETODHVD
BCoV nsn2	(204)	ENCOUNTY SE	TANK COL	- tip	WELL TIME	EDFIGURACIA Y	GESONKEDL
Consensus	(205)	VVAWLYAA:	TN CN	5177	TOCOMOO!	P.D.R. L. A. W. WESSER	CK SOMKEDF
	(200)	VVANDIAA.	LIN CN	ex E	POSDICSE	EDFHUWAMSN	
	(256)	256	27	^	000	000	Section 6
swinn IRV nen2	(200)	a Horse Wallet	21	 	280	290	DELTPESTENO
MEN pen?	(251)	THE TANK	COVDVCKI	LRTIMV	KNS-Q觸GG	DPILGOYNE	DELTPESTERQ
SAPS non2	(241)	VIDALASM	CAMALON	LAAIKR	LHS-GFQC	CILCISCACE	DELTPSDYYQQ
PCoV non2	(243)	THE PLEAD	CHAVLDE	CAANKE	Pronguido.	RTILCSTATE	DEFTEFOVVRQ
BCOV risp2	(247)	ATIDA LASM.	VSLETI	LAAIKR	LKN-GFQG	RQIMOSCSFE	DELTESDVYQQ
Consensus	(256)	VIDALAAM:	GVSVE I	LAAIKR	L S GFQG	RQILGSCILE	DELTPSDVYQQ
							Section 7
	(307)						
avian IBV nsp2	(301)	TECABLO		NO: 6570			
		PYCAKPO		NO: 6571			
SARS nsp2				NO: 6569			
BCoV nsp2	(297)	LAGEKLO		NO: 6572			i
Consensus	(307)	LAGVKLQ	SEQ ID	NO: 6573			1

FIGURE 17

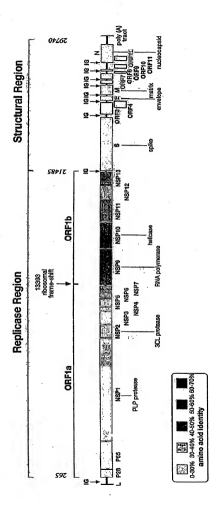
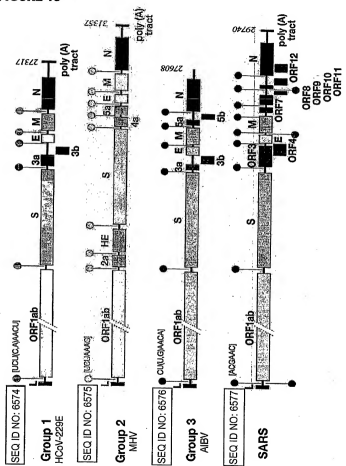
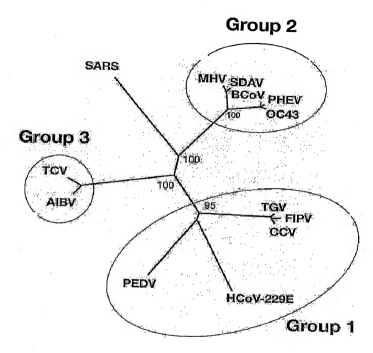


FIGURE 18



FUNCTION STRUCTURE Receptor binding (globular domain) Oligomerization domain Leucine zipper (stalk domain) Oligomerization domain Leucine Plasma zipper membrane Membrane anchor Hydrophobic domain Cysteine-rich domain Cytoplasmic tail



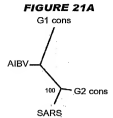
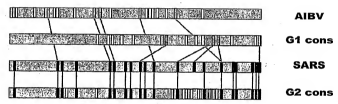
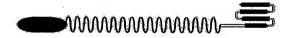
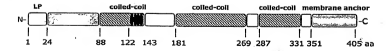


FIGURE 21B







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FIGURE 23

LPRKSQPTSISCRSVL-TNFKICVAVARLHA-CTYAV-TIINFTVVDKKRVTRPSSADCL RFRPCCSRSSAYLGFVRV-PKGKMESLVLGVNEKTHVQLSLPVLQVRDVLVRGFGDSVEE ALSEAREHLKNGTCGLVELEKGVLPOLEQPYVFIKRSDALSTNHGHKVVELVAEMDGIOY GRSGITLGVLVPHVGETPIAYRNVLLRKNGNKGAGGHSYGIDLKSYDLGDELGTDPIEDY EONWNTKHGSGALRELTRELNGGAVTRYVDNNFCGPDGYPLDCIKDFLARAGKSMCTLSE OLDYIESKRGVYCCRDHEHEIAWFTERSDKSYEHOTPFEIKSAKKFDTFKGECPKFVFPL NSKVKVIQPRVEKKKTEGFMGRIRSVYPVASPQECNNMHLSTLMKCNHCDEVSWQTCDFL KATCEHCGTENLVIEGPTTCGYLPTNAVVKMPCPACODPEIGPEHSVADYHNHSNIETRL RKGGRTRCFGGCVFAYVGCYNKRAYWVPRASADIGSGHTGITGDNVETLNEDLLEILSRE RVNINIVGDFHLNEEVAIILASFSASTSAFIDTIKSLDYKSFKTIVESCGNYKVTKGKPV KGAWNIGQQRSVLTPLCGFPSQAAGVIRSIFARTLDAANHSIPDLQRAAVTILDGISEOS LRLVDAMVYTSDLLTNSVIIMAYVTGGLVQQTSOWLSNLLGTTVEKLRPIFEWIEAKLSA GVEFLKDAWEILKFLITGVFDIVKGQIQVASDNIKDCVKCFIDVVNKALEMCIDQVTIAG AKLRSLNLGEVFIAQSKGLYRQCIRGKEQLQLLMPLKAPKEVTFLEGDSHDTVLTSEEVV LKNGELEALETPVDSFTNGAIVGTPVCVNGLMLLEIKDKEOYCALSPGLLATNNVFRLKG GAPIKGVTFGEDTVWEVQGYKNVRITFELDERVDKVLNEKCSVYTVESGTEVTEFACVVA ${\tt EAVVKTLQPVSDLLTNMGIDLDEWSVATFYLFDDAGEENFSSRMYCSFYPPDEEEEDDAE}$ CEEEEIDETCEHEYGTEDDYQGLPLEFGASAETVRVEEEEEEEDWLDDTTEQSEIEPEPEP TPEEPVNQFTGYLKLTDNVAIKCVDIVKEAQSANPMVIVNAANIHLKHGGGVAGALNKAT NGAMQKESDDYIKLNGPLTVGGSCLLSGHNLAKKCLHVVGPNLNAGEDIQLLKAAYENFN SQDILLAPLLSAGIFGAKPLQSLQVCVQTVRTQVYIAVNDKALYEQVVMDYLDNLKPRVE APKQEEPPNTEDSKTEEKSVVQKPVDVKPKIKACIDEVTTTLEETKFLTNKLLLFADING KLYHDSQNMLRGEDMSFLEKDAPYMVGDVITSGDITCVVIPSKKAGGTTEMLSRALKKVP VDEYITTYPGQGCAGYTLEEAKTALKKCKSAFYVLPSEAPNAKEEILGTVSWNLREMLAH AEETRKLMPICMDVRAIMATIORKYKGIKIOEGIVDYGVRFFFYTSKEPVASIITKLNSL NEPLVTMPIGYVTHGFNLEEAARCMRSLKAPAVVSVSSPDAVTTYNGYLTSSSKTSEEHF VETVSLAGSYRDWSYSGQRTELGVEFLKRGDKIVYHTLESPVEFHLDGEVLSLDKLKSLL ${\tt SLREVKTIKVFTTVDNTNLHTQLVDMSMTYGOOFGPTYLDGADVTKIKPHVNHEGKTFFV}$ LPSDDTLRSEAFEYYHTLDESFLGRYMSALNHTKKWKFPQVGGLTSIKWADNNCYLSSVL LALQQLEVKFNAPALQEAYYRARAGDAANFCALILAYSNKTVGELGDVRETMTHLLQHAN LESAKRVLNVVCKHCGQKTTTLTGVEAVMYMGTLSYDNLKTGVSIPCVCGRDATQYLVQQ ESSFVMMSAPPAEYKLQQGTFLCANEYTGNYQCGHYTHITAKETLYRIDGAHLTKMSEYK GPVTDVFYKETSYTTTIKPVSYKLDGVTYTEIEPKLDGYYKKDNAYYTEQPIDLVPTQPL ${\tt PNASFDNFKLTCSNTKFADDLNQMTGFTKPASRELSVTFFPDLNGDVVAIDYRHYSASFK}$ KGAKLLHKPIVWHINQATTKTTFKPNTWCLRCLWSTKPVDTSNSFEVLAVEDTQGMDNLA CESQQPTSEEVVENPTIQKEVIECDVKTTEVVGNVILKPSDEGVKVTQELGHEDLMAAYV ENTSITIKKPNELSLALGLKTIATHGIAAINSVPWSKILAYVKPFLGQAAITTSNCAKRL AQRVFNNYMPYVFTLLFQLCTFTKSTNSRIRASLPTTIAKNSVKSVAKLCLDAGINYVKS PKFSKLFTIAMWLLLLSICLGSLICVTAAFGVLLSNFGAPSYCNGVRELYLNSSNVTTMD FCEGSFPCSICLSGLDSLDSYPALETIQVTISSYKLDLTILGLAAEWVLAYMLFTKFFYL LGLSAIMQVFFGYFASHFISNSWLMWFIISIVQMAPVSAMVRMYIFFASFYYIWKSYVHI ${\tt MDGCTSSTCMMCYKRNRATRVECTTIVNGMKRSFYVYANGGRGFCKTHNWNCLNCDTFCT}$ GSTFISDEVARDLSLQFKRPINPTDQSSYIVDSVAVKNGALHLYFDKAGQKTYERHPLSH FVNLDNLRANNTKGSLPINVIVFDGKSKCDESASKSASVYYSQLMCQPILLLDQALVSDV ${\tt GDSTEVSVKMFDAYVDTFSATFSVPMEKLKALVATAHSELAKGVALDGVLSTFVSAARQG}$ VVDTDVDTKDVIECLKLSHHSDLEVTGDSCNNFMLTYNKVENMTPRDLGACIDCNARHIN AQVAKSHNVSLIWNVKDYMSLSEQLRKQIRSAAKKNNIPFRLTCATTRQVVNVITTKISL KGGKIVSTCFKLMLKATLLCVLAALVCYIVMPVHTLSIHDGYTNEIIGYKAIQDGVTRDI ISTDDCFANKHAGFDAWFSQRGGSYKNDKSCPVVAAIITREIGFIVPGLPGTVLRAINGD FLHFLPRVFSAVGNICYTPSKLIEYSDFATSACVLAAECTIFKDAMGKPVPYCYDTNLLE GSISYSELRPDTRYVLMDGSIIQFPNTYLEGSVRVVTTFDAEYCRHGTCERSEVGICLST SGRWVLNNEHYRALSGVFCGVDAMNLIANIFTPLVQPVGALDVSASVVAGGIIAILVTCA AYYFMKFRRVFGEYNHVVAANALLFLMSFTILCLVPAYSFLPGVYSVFYLYLTFYFTNDV SFLAHLQWFAMFSPIVPFWITAIYVFCISLKHCHWFFNNYLRKRVMFNGVTFSTFEEAAL

CTFLLNKEMYLKLRSETLLPLTOYNRYLALYNKYKYFSGALDTTSYREAACCHLAKAI.ND FSNSGADVLYOPPOTSITSAVLOSGFRKMAFPSGKVEGCMVOVTCGTTTLINGLWLDDTVY CPRHVICTAEDMLNPNYEDLLIRKSNHSFLVOAGNVQLRVIGHSMQNCLLRLKVDTSNPK TPKYKFVRIQPGQTFSVLACYNGSPSGVYQCAMRPNHTIKGSFLNGSCGSVGFNIDYDCV SFCYMHHMELPTGVHAGTDLEGKFYGPFVDROTAOAAGTDTTITLNVLAWLYAAVINGDR WFLNRFTTTLNDFNLVAMKYNYEPLTQDHVDILGPLSAOTGIAVLDMCAALKELLONGMN GRTILGSTILEDEFTPFDVVROCSGVTFQGKFKKIVKGTHHWMLLTFLTSLLILVOSTOW SLFFFVYENAFLPFTLGIMAIAACAMLLVKHKHAFLCLFLLPSLATVAYFNMVYMPASWV MRIMTWLELADTSLSGYRLKDCVMYASALVLLILMTARTVYDDAARRVWTI,MNYITI,VYK VYYGNALDOAISMWALVISVTSNYSGVVTTIMFLARAIVFVCVEYYPLLFITGNTLOCIM LVYCFLGYCCCCYFGLFCLLNRYFRLTLGVYDYLVSTOEFRYMNSOGLLPPKSSIDAFKL NIKLLGIGGKPCIKVATVQSKMSDVKCTSVVLLSVLQQLRVESSSKLWAQCVQLHNDILL AKDTTEAFEKMVSLLSVLLSMQGAVDINRLCEEMLDNRATLQAIASEFSSLPSYAAYATA OEAYEOAVANGDSEVVLKKLKKSLNVAKSEFDRDAAMQRKLEKMADQAMTOMYKOARSED KRAKVTSAMOTMLFTMLRKLDNDALNNIINNARDGCVPLNIIPLTTAAKLMVVVPDYGTY KNTCDGNTFTYASALWEIQQVVDADSKIVQLSEINMDNSPNLAWPLIVTALRANSAVKLO NNELSPVALROMSCAAGTTQTACTDDNALAYYNNSKGGRFVLALLSDHODLKWARFPKSD GTGTIYTELEPPCRFVTDTPKGPKVKYLYFIKGLNNLNRGMVLGSLAATVRLOAGNATEV PANSTVLSFCAFAVDPAKAYKDYLASGGQPITNCVKMLCTHTGTGQAITVTPEANMDOES FGGASCCLYCRCHIDHPNPKGFCDLKGKYVOIPTTCANDPVGFTLRNTVCTVCGMWKGYG CSCDOLREPLMQSADASTFLNGFAV-VQPVLHRAAQALVLMSSTGLLIFTTKKLLVLQSS -KLIAVASRRRMRKAIY-TLTL-LRGILCLTTNMKRLFITWLKIVORLLSMTFSSLE-MV TWYHIYHVSV-LNTOWLI-SMLYVILMRVIVIH-KKYSSHTIAVMMIISIRRIGMTS-RI LTSYAYMLT-VSVYANHY-RLYNSAMLCVMQAL-AY-H-IIRILMGTGTISVISYK-HOA AEFLLWIHITHC-CPSSL-LGHWLLSPIWMLISONHLLSGIC-NMIIRKRDFVSSTVIIN IGTRHTIPIVLTVWMIGVSFIVQTLMCYFLLCFHLOVLDH--EKYL-MVFLLLFOLDTIF VS-ESYIIRM-TYIARVSVSRNF-CMLLIQLCMQLLAIYC-INALHAFQ-LH-QTMLLFK LSNPVILIKTFMTLLCLKVSLRKEVLLN-NTSSLLRMATLLSVIMTIIVIICQQCVISDN SYS-LKLLINTLIVTMVAVLMPTK-SLTIWINOLVSHLINGVRLDFIMTO-VMRIKMHFS RILSVMSSLL-LK-ILSMPLVQRIELAP-LVSLSVVL-QIDSFIRNY-SO-PPLEELLW-LEQASFTVAGIIC-KLFTVM-KLHTLWVGIIQNVTEPCLTCLG-WPLLFLLANITLAVTY HTVSTG-LTSVRKY-VRWSCVAAHYMLNOVEHHPVMLOLLMLIVSLTFVKLLOPM-MHFF QLMVIR-LTSMSAIYNTGSMSVSIEIGMLIMNSWMSFTLTCVNISP--FFLMMPLCAITV TMRLKV--LALRTLRQFFIIKIMCSCLRQNVGLRLTLLKDLTNFAHSIOC-LNKEMITCT CLTQIHQEY-AQAVLSMILSKQMVHL-LKGSCHWLLMLTHLQNILIRSMLMSFTCIYNTL ESYMMSLLATCWTCIP-C-LMITPHGTGNLSFMRLCTHHIQSCRL-VLVYCAIHRLHFVA VPVLGDHSYVASAAMTMSFQHHTN-CCLLIPMFAMPQVVMSLM-HNCI-EV-AIIASHIS LPLVFHYVLMVRFLVYTKTHV-AVTMSLTSMR-QHVIGLMLAITYLPTLVLRDSSFSQQK RSKPLRKHLSCHMVLPLYAKYSLTENCIFHGRLENLDHH-TETMSLLVTV-LKIVKYRLE STPLKKVTMVMLLCTEVLRHTS-MLVITLC-HLTL-CHLVHLL-CHKSTM-ELLACTOHS TSOMSFLAMLOIIKRSACKSTLHSKDHLVLVRVILPSDLLSITHLLA-CTRHAIMOLLMP YVKRH-NICP-INVVESYLRVRA-SVLINSK-IQH-NSMFSAL-MHCQKQLLTL-SLMKS LWLLIMT-VLSMLDFVQNTTSILAILLNYQPPAHC-LKAH-NONILIOCADL-KO-VOTC SLELVAVVLLKLLTL-VL-FMTIS-KHTRISOLNASKCSTKVLLHMMFHIOSTDIK-AI,-ENFLHAILLGEKLFLSHLIIHRTL-LQKS-DCLRRLLIHHRVLNMTMSYSHKLLKOHTLV MSTASMWLSQGQKLAFCA-CLIEIFMTNCNLQV-KYHVAMWLHYKQKM-LDFLRTVVRSL LVFILHRHLHTSALI-SSRLKDYVLTYQAYQRT-PTVDSSL-WVSK-ITKSMVTLICLSP AKKLFVTFVRGLALM-RAVMQLEMLWVLTYLSS-DFLQVLT--LYRLVMLTLKITONSPE LMQNLHQVTSLNILYHSCIKACPGM-CVLR-YKCSVIH-KDCOTESCSSFGRMALSLHO-STLSRLDLKERVVCVTNVQLAFLLHQILMPAGIILWVLTMSITHL-LMFSSGALRVTFRV TMTNIARYMEMHMWLVVMLS-LDV-QSMSALLSALIGLLNTLL-EMN-GLILLAEKYNTW L-SLHCLLISFQFFMTLEIQRLSSVCLRLK-NGSSTMLSHVVTKLTK-RNSSILMLHITI NSLMVFVCFGIVTLIVTQPMQLCVGLTQESCQT-TYQAVMVVVCM-ISMHSTLQLSIKVH LLI-SNCLSFTILIVLVSLMANK-CRILIMFHSNLLRVLHDAI-VVLFADTMOMSTDSTW MHII--FLLDLAYGFTNNLILITCGIHLPGYRV-KMWLIMLLIKDTLMDTPAKHLFPSLI MLFTOR-MVLMWRSLKIROHFLLMLHLSFGLSVTLNOCORLRYSIIWVLISLLIL-SGTT

KEKPOHMYLO-VSAO-LTLPRNLLRVLVLHLLSCLMVEWKDR-TFLETPVMVF--OKVOS KV-HLQRDQHKLASMESH-LENQ-KHSLTTLRK-TALFNSCLKPTLLRAET-RILSPDHK WKLTFSSSLWMNSYSDISSRAMPSNTSFMEISVMDNLAVFI---A-PSAHKIHHLN-RIL SLWTAQ-KITS-OMRKOVHONVCVL-LIFYLMTLSR--SHKICQ-FOKWSRLQLTMLKFH SCFGVRMDMLKPSTONYKOVERGNOVLRCLTCTRCKECFLKSVTFRIMVKMLLYOKE---MSOSILNCVNT-IHLL-LYPTT-ELFTLVLALIKELHQVOLCSDNGCOLAHYLSIOILMT SSPTHILL-LETVOOYIRLINGTLLLAICMTLGPNM-QKRMTLKKGFSLICVDL-SKN-P WVVL-L-R-OSILGMLTFTSLWAISHGGQLLLQM-MHHRKHF-LGLTILASRRNKLMAI PCMLTTFSGGTQILSSCLPIHSLT-ANFLLN-EELL-CLLRRIKSMI-FILFWKKVGLSL EKTTELWFQVIFLLTTKRTCLFSYYFLLSLVVVTLTGAPLLMMFKLLITLNILHL-GGFT ILMKFLDQTLFI-LRIYFFHFILMLQGFILLIIRLATLSYLLRMVFILLPQRNQMLSVVG FLVLP-TTSHSR-LLLTILLMLLYEHVTLNCVTTLSLLFLNPWVHRHIL-YSIMHLIALS STYLMPFRLMFQKSQVILNTYESLCLKIKMGFSMFIRAINL-M-FVIYLLVLTL-NLFLS CLLVLTLQILEPFLQPFHLLKTFGARQLQPILLAI-SQLHLCSSMMKMVQSQMLLIVLKI HLLNSNALLRALRLTKEFTRPLISGLFPQEML-DSLILOTCVLLERFLMLLNSLLSMHGR EKKFLIVLLITLCSTTQHFFQPLSAMAFLPLS-MIFASPMSMQILL-SREMM-DK-RODK LVLLLIIIINCQMISWVVSLLGILGTLMLLQLVIIIINIGILDMASLGPLRETYLMCLSP LMANLAPHLLLIVIGH-MIMVFTPLLALATNLTEL-YFLLNF-MHRPRFVDQNYPLTLLR TSVSILILMDSLVLVC-LLLQRDFNHFNNLAVMFLISLIPFEILKHLKY-TFHLALLGV-V-LHLEQMLHLKLLFYIKMLTALMFLOQFMOINSHOLGAYILLETMYSRLKOAVL-ELSM STLLMSATFLLELAFVLVTIQFLYYVVLAKNLLWLILCL-VLIVQLLTLITPLLYLLTFO LALLQK-CLFLWLKPP-IVICTSAEILLNVLICFSNMVAFAHN-IVHSQVLLLNRIATHV KCSLKSNKCTKPQL-NILVVLIFHKYYLTL-SQLRGLLLRTCSLIR-HSLMLAS-SNMAN A-VILMLEISFVRRSSMDLQCCHLCSLMI-LLPTLLL-LVVLPLLDGHLVLALLFKYLLL CKWHIGSMALELPKMFSMRTKNKSPTNLTRRLVKFKNHLQQHQLHWASCKTLLTRMLKH-THLLNNLALILVQFQVC-MISFRDLIKSRRRYKLTG-LQADFKAFKPM-HNN-SGLLKSG LLLILLLKCLSVFLDNQKELTFVERATTLCPSHKQPRMVLSSYMSRMCHPRRGTSPORO QFVMKAKHTSLVKVFLCLMALLGLLHRGTSFLHK-LLQTIHLSQEIVMSLLASLTTQFMI LCNLSLTHSKKSWTSTSKIIHHQMLILATFQALTLLSSTFKKKLTASMRSLKI-MNHSLT FKNWENMSNILNGLGMFGSASLLD-LPSSWLOSCFVA-LVVAVASRVHALVVLAASLMRM TLSQFSRVSNYITHKRTYGFVYEIFYSWINYCTASKN-QCFSCKYCSCYSNDTATSLTPF RMACYWRCISCCFSERYQNNCAQ-KMAASPL-GLPVHLQFTAAICYHLFTSFACRCRYGG AIFVPLCLDIFSTMHQRM-NYYEMLALLEVQIQEPITL-CQLLCLLAHT-L-LLYTI-OC HRYNCRY-R-RHFNTKTQRRLPNWWLF-G-ALRC-RLCRCTWLFHRSLLPA-VYTNYYRH WY-KCYILHL-QAC-RPTECANTHNRRLFRSC-SSNGSNL--ADDDY-RAFVSTRK-VRT YVLIRFGRNRYVNS--RTSFSCFRGILASHTSHPYCASIVCVLLQYC-REFSKTNGLRLL AC-KSELF-RSS-SSGLNELTIIIILFGTLTLLIMADNGTITVEELKOLLEQWNLVIGFL FLAWIMLLQFAYSNRNRFLYIIKLVFLWLLWPVTLACFVLAAVYRINWVTGGIAIAMACT VGLMWLSYFVASFRLFARTRSMWSFNPETNILLNVPLRGTIVTRPLMESELVIGAVIIRG HLRMAGHSLGRCDIKDLPKEITVATSRTLSYYKLGASQRVGTDSGFAAYNRYRIGNYKLN TDHAGSNDNIALLVQ-VTTDVSSC-LPGYNSRDIDYHYEDFQDCYLES-RYNKFNSETII -ASN-EELFGVR--RTYGVRLSIKRT-KLFSS-H-LYLHLASYITIRSVLEVRLYY-KNL $\mathtt{AHQEHTRAIHHFTLLLTINLH-LALAHTLLLLVLTVLDIPISCVQDQFHQNFSSDKRRFN}$ KSSTRHFFSLLLL-YF-YFASPLRERQNE-AHFN-LLFVLFSLSAIPCFNNAYYILVFTR NPGSRRTLYQSLNEHETSHCFDLYFSMQLHMHCSTALCI--TSCA-RSL-GTTLGVILIA LLGFVL-ERFYLFIDGTLWFKHAHLMLLSTVKIQLVVRL-LGVGTFMKVTKLLHLETYLL F-INEQIKMSDNGPQSNQRSAPRITFGGPTDSTDNNQNGGRNGARPKQRRPQGLPNNTAS WFTALTOHGKEELRFPRGQGVPINTNSGPDDQIGYYRRATRRVRGGDGKMKELSPRWYFY YLGTGPEASLPYGANKEGIVWVATEGALNTPKDHIGTRNPNNNAATVLOLPOGTTLPKGF YAEGSRGGSQASSRSSSRSRGNSRNSTPGSSRGNSPARMASGGGETALALLLLDRLNOLE ${\tt SKVSGKGQQQQGQTVTKKSAAEASKKPRQKRTATKQYNVTQAFGRRGPEQTQGNFGDODL}$ IROGTDYKHWPQIAOFAPSASAFFGMSRIGMEVTPSGTWLTYHGAIKLDDKDPQFKDNVI LLNKHIDAYKTFPPTEPKKDKKKKTDEAQPLPQRQKKQPTVTLLPAADMDDFSRQLQNSM SGASADSTQA-TLMMTTQGRWAM-TFSQFRLRYIVYSCAE-ILVTKQHK-V-LTLISHSN L-SMCNIREDLKEPPHFHRGHAEYDRGYSE-C-GELPIWKSPNV-N-F-CYPHVILIAS

TOEKPTNLDLL-ICSLNEL-NLCSCRSAACLVHLRSINNNKFYCR-OETSNSSLFCRLLT VSSVLQSIISIPRFRPGVTER-DGEPCSWCQRENTRPTQFACPSG-RRASAWLRGLCGRG PIGGT-TPOKWHLWSSRAGKRRTAPA-TALCVH-TF-CLKHOSRPOGR-AGCRNGRHSVR S-RYNTGSTRATCGRNPNCIPQCSSS-ER--GSRWS-LWHRSKVL-LR-RAWH-SH-RL-TKLEH-AWOWCTP-THS-AOWRCSHSLCROOFLWPRWVPS-LHORFSRTRGOVNVHSFRT T-LHRVEERCLLLP-P-A-NCLVH-AL--ELRAPDTLRN-ECQEI-HFORGMPKVCVSS-LKSOSHSTTC-KEKD-GFHGAYTLCVPCCISTGV-OYALVYLDEM-SLR-SFMADVRISE SHL-TLWH-KFSY-RTYYMWVPTY-CCSENAMSCLSRPRDWT-A-CCRLSQPLKH-NSTP QGR-D-MFWRLCVCLCWLL--ACLLGSSC-C-YWLRPYWHYW-OCGDLE-GSP-DTES-T C-H-HCWRFSFE-RGCHHFGIFLCFYKCLY-HYKES-LQVFQNHC-VLR-L-SYQGKARK RCLEHWTTEISFNTTVWFSLTGCWCYOINFCAHT-CSKPLNS-FAKSSCHHT-WYF-TVI TSCRRHGLYFRPAHOQCHYYGICNWWSCTTDFSVVV-SFGHYC-KTOAYL-MD-GET-CR S-ISQGCLGDSQISHYRCF-HRQGSNTGCFR-HQGLCKMLH-CC-QGTRNVH-SSHYRWR KVAITOLR-SLHRSKQGTLPSVYTWQGAAATTHAS-GTKRSNLS-R-FT-HSTYL-GGCS QER-TRSTRDAR--LHKWSYRRHTSLCKWPHALRD-GQRTILRIVSWFTGYKOCLSLKRG CTN-RCNLWRRYCLGSSRLOECENHI-A--TC-OSA--KVLCLHC-IRYRSY-VCMCCSR GCCEDFTTSF-SPYQHGY-S--VECSYILLI--CW-RKLFITYVLFLLPSR-GRRGRCRV -GRRN--NL-T-VRYRG-LSRSPSGIWCLS-NSSS-GRRRGRLAG-YY-AIRD-ARTRTY T-RTS-SVYWLFKTY-OCCH-MC-HR-GGTKC-SYGDCKCC-HTPETWWWCSRCTOOGNO WCHAKGE--LH-AKWPSYSRRVLFAFWT-SC-EVSACCWT-PKCR-GHPAS-GSI-KFOF TGHLTCTIVVSRHIWC-TTSVFTSVRADGSYTGLYCSQ-QSSL-AGCHGLS--PEA-SGS T-TRGATKHRRFQN-GEICRTEACRCEAKN-GLH--GYHNTGRN-VSYO-VTLVC-YOW-ALP-FSEHA-R-RYVFP-EGCTLHGR-CYH-W-YHLCCNTLOKGWWHY-DALKSFEESAS --VYNHVPWTRMCWLYT-GS-DCS-EMQICILCTTFRST-C-GRDSRNCILEFERNACSC -RDKKINAYMHGC-SHNGNHPT-V-RN-NSRGHR-LWCPILLLY--RACSFYYYEAELSK -AACHNANWLCDTWF-S-RGCALYAFS-SSCRSVSIITRCCYYI-WIPHFVIKDI-GALC RNSFFGWLLQRLVLFRTAYRVRC-IS-AW-ONCVPHSGEPRRVSS-R-GSFT-OTKESLI PAGG-DYKSVHNCGQH-SPHTACGYVYDIWTAVWSNILGWC-CYKN-TSCKS-G-DFLCT T---HTT--SFRVLPYS--EFSW-VHVCFKPHKEMEISSSWWFNFN-MG--OLLFV-CFI STSTA-SQIQCTSTSRGLL-SPCW-CC-LLCTHTRLQ--NCWRAW-CORNYDPSSTAC-F GICKASS-CGV-TLWSENYYLNGCRSCDVYGYSIL--S-DRCFHSMCVWS-CYTISSTTR VFFCYDVCTTC-V-ITARYILMCE-VHW-LSVWSLHSYNC-GDPLSY-RSSPYKDVRVOR TSD-CFLQGNILHYNHOACVV-TRWSYLHRD-TKIGWVL-KG-CLLYRAAYRPCTNSTIT KCEF--FQTHMF-HKIC--FKSNDRLHKASFTRAICHILPRLEWRCSGY-L-TLFSEFOE RC-ITA-ANCLAH-PGYNQDNVQTKHLVFTLSLEYKASRYFKFI-SSGSRRHTRNGOSCL -KSTTHL-RSSGKSYHTEGSHRV-RENYRSCRQCHT-TIR-RC-SNTRVRS-GSYGCLCG KHKHYH-ET--AFTSLRFKNNCHSWYCCN--CSLE-NFGLCQTILRTSSNYNIKLR-EIS TTCV-QLYALCVYIIVPIVYFY-KYOF-N-SFTTYNYC-K-C-ECC-IMFGCRH-LCEVT OIF-IVHNRYVAIVVKYLLRFSNLCNCCFWCTLI-FWCSFLL-WR-RTVS-FV-RYYYGF L-RFFSLQHLFKWIRLP-FLSSS-NHSGDDFIVQARLDNFRSGR-VGFGIYVVHKILLFI RSFSYNAGVLWLFC-SFHQQFLAHVVYH-YCTNGTRFCNG-DVHLLCFFLLHMEELCSYH GWLHLFDLHDVL-AOSCHTR-VYNYC-WHEEIFLCLCKWRPWLLODSOLELSOL-HILHW -YIH---SCS-FVTPV-KTNOPY-PVIVYC--CCCEKWRASPLL-QGWSKDL-ETSALPF CQFRQFES-QH-RFTAY-CHSF-WQVQMRRVCF-VCFCVLQSADVPTYSVA-PSSCIRRW R-Y-SFR-DV-CLCRHLFSNF-CSYGKT-GTCCYSSORVSKGCSFRWCPFYIRVSCPTRC C-YRC-HKGCY-MSQTFTSL-LRSDR-QL-OFHAHL--G-KHDAORSWRMY-L-CKAYOC PSSKKSQCFTHLECKRLHVFI-TAA-TNS-CCQEEQHTF-TNLCYN-TGCQCHNY-NLTQ GW-DC-YLF-TYA-GHIIVRSCCIGLLYRYASTYIVNP-WLHK-NHWLOSHSGWCHS-HH FY--LFCK-TCWF-RMV-PAWWFIQK-QKLPCSSCYHYKRDWFHSAWLTGYCAESNOW-L LAFSTSCF-CCWQHLLHTFQTH-V--FCYLCLRSCC-VYNF-GCYGOTCAILL-H-FARG FYFL--ASSRHSLCAYGWFHHTVS-HLPGGFC-SSNNF-C-VL-TWYMRKVRSRYLPIYO W-MGS---ALQSSIRSFLWC-CDESHS-HLYSSCATCGCFRCVCFSSGWWYYCHIGDLCC LLLYEIQTCFW-VQPCCCC-CTFVFDVFHYTLSGTSLOLSAGSLLSLLLVLDILFHO-CF ILGSPSMVCHVFSYCAFLDNSNLCILYFSEALPLVL-QLS-EKSHV-WSYI-YLRGGCFV

YLFAOQGNVPKIA-RDTVATYTV-OVSCSI-QVQVFQWSLRYYQLS-SSLLPLSKGSK-L -OLRC-CSLPTTTDINHFCCSAEWF-ENGIPVROS-RVHGTSNLWNYNS-WIVVG-HSTI SKTCHLHSRRHA-S-L-RSAHSQIOP-LSCSGWQCSTSCYWPFYAKLSA-A-S-YF-P-D TQV-ICPYPTWSNIFSSSMLOWFTIWCLSVCHET-SYH-RFFP-WIMW-CWF-H-L-LRV FLLYASYGASNRSTRWY-LRR-ILWSIC-QTNCTGCRYRHNHNIKCFGMAVCCCYOW--V VS--IHHYFE-L-PCGNEVOL-TFDTRSC-HIGTSFCSNRNCRLRYVCCFERAAAEWYEW SYYPW-HYFRR-VYTI-CC-TMLWCYLPR-VOENC-GHSSLDAFNFLDITIDSCSKYTVV TVFLCLRECFLAIYSWYYGNCCMCYAAC-A-ARILVLVSVTFSCNSCLL-YGLHAC-LGD AYHDMA-IG-H-LVWL-A-GLCYVCFSFSFAYSHDSSHCL--CC-TCLDTDECHYTCLOS LLW-CFRSSYFHVGLSYFCNL-LFWCRYDYHVFS-SYSVCVC-VLPIVIYYWOHLTVYHA CLLFLRLLLLLLWPFLFTQPLLQAYSWCL-LLGLYTRI-VYELPGAFAS-E-Y-CFOA-H-VVGYWR-TMYQGCYCTV-NV-RKVHICGTALGSSTT-SRVIF-IVGTMCTTPQ-YSSC KRHN-SFREDGFSFVCFAIHAGCCRH--VVRGNAR-PCYSSGYCFRI-FFTIICRLCHCP GGL-AGCS-W-F-SRSQKVKEIFECG-I-V-P-CCHATOVGKDGRSGYDPNVOTGKI-GO EGKSN-CYANNALHYA-EA---CT-QHYOOCA-WLCSTQHHTIDYSSQTHGCCP-LWYLO EHL-W-HLYICICTLGNPASC-CG-QDCST--N-HGQFTKFGLASYCYSSKSQLSC-TTE --TESSSTTTDVLCGWYHTNSLY--QCTCLL-QFEGR-VCAGITIRPPRSQMG-IP-E-W YRYNLHRTGTTL-VCYRHTKRA-SEILVLHQRLKQPK-RYGAGQFSCYSTSSGWKCYRST COFNCAFLLCFCSRPC-SI-GLPSKWRTTNHOLCEDVVYTHWYRTGNYCNTRS-HGPRVL WWCFMLSVL-MPH-PSKS-RIL-LER-VRPNTYHLC--PSGFYT-KHSLYRLRNVERLWL -L-PTPRTLDAVCGCINVFKRVCGVSAARLTPCGTGTSTDVVYRAFDIYNEKVAGFAKFL KTNCCRFQEKDEEGNLLDSYFVVKRHTMSNYQHEETIYNLVKDCPAVAVHDFFKFRVDGD MVPHISRQRLTKYTMADLVYALRHFDEGNCDTLKEILVTYNCCDDDYFNKKDWYDFVENP DILRVYANLGERVROSLLKTVOFCDAMRDAGIVGVLTLDNODLNGNWYDFGDFVOVAPGC GVPIVDSYYSLLMPILTLTRALAAESHMDADLAKPLIKWDLLKYDFTEERLCLFDRYFKY WDQTYHPNCINCLDDRCILHCANFNVLFSTVFPPTSFGPLVRKIFVDGVPFVVSTGYHFR ELGVVHNQDVNLHSSRLSFKELLVYAADPAMHAASGNLLLDKRTTCFSVAALTNNVAFOT VKPGNFNKDFYDFAVSKGFFKEGSSVELKHFFFAQDGNAAISDYDYYRYNLPTMCDIROL LFVVEVVDKYFDCYDGGCINANQVIVNNLDKSAGFPFNKWGKARLYYDSMSYEDODALFA YTKRNVIPTITQMNLKYAISAKNRARTVAGVSICSTMTNRQFHQKLLKSIAATRGATVVI GTSKFYGGWHNMLKTVYSDVETPHLMGWDYPKCDRAMPNMLRIMASLVLARKHNTCCNLS HRFYRLANECAQVLSEMVMCGGSLYVKPGGTSSGDATTAYANSVFNICQAVTANVNALLS TDGNKIADKYVRNLQHRLYECLYRNRDVDHEFVDEFYAYLRKHFSMMILSDDAVVCYNSN YAAQGLVASIKNFKAVLYYQNNVFMSEAKCWTETDLTKGPHEFCSQHTMLVKQGDDYVYL PYPDPSRILGAGCFVDDIVKTDGTLMIERFVSLAIDAYPLTKHPNQEYADVFHLYLOYIR KLHDELTGHMLDMYSVMLTNDNTSRYWEPEFYEAMYTPHTVLQAVGACVLCNSOTSLRCG ACIRRPFLCCKCCYDHVISTSHKLVLSVNPYVCNAPGCDVTDVTOLYLGGMSYYCKSHKP PISFPLCANGQVFGLYKNTCVGSDNVTDFNAIATCDWTNAGDYILANTCTERLKLFAAET LKATEETFKLSYGIATVREVLSDRELHLSWEVGKPRPPLNRNYVFTGYRVTKNSKVOIGE YTFEKGDYGDAVVYRGTTTYKLNVGDYFVLTSHTVMPLSAPTLVPQEHYVRITGLYPTLN ISDEFSSNVANYOKVGMOKYSTLOGPPGTGKSHFAIGLALYYPSARIVYTACSHAAVDAL CEKALKYLPIDKCSRIIPARARVECFDKFKVNSTLEQYVFCTVNALPETTADIVVFDEIS MATNYDLSVVNARLRAKHYVYIGDPAQLPAPRTLLTKGTLEPEYFNSVCRLMKTIGPDMF LGTCRRCPAEIVDTVSALVYDNKLKAHKDKSAQCFKMFYKGVITHDVSSAINRPQIGVVR EFLTRNPAWRKAVFISPYNSQNAVASKILGLPTQTVDSSQGSEYDYVIFTQTTETAHSCN VNRFNVAITRAKIGILCIMSDRDLYDKLQFTSLEIPRRNVATLQAENVTGLFKDCSKIIT GLHPTQAPTHLSVDIKFKTEGLCVDIPGIPKDMTYRRLISMMGFKMNYQVNGYPNMFITR EEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQLGFSTGVNLVAVPTGYVDTENNTEFTRV NAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQMLSDTLKGLSDRVVFVLWAHGFELTSMK YFVKIGPERTCCLCDKRATCFSTSSDTYACWNHSVGFDYVYNPFMIDVQQWGFTGNLOSN $\verb|HDQHCQVHGNAHVASCDAIMTRCLAVHECFVKRVDWSVEYPIIGDELRVNSACRKVQHMV|\\$ VKSALLADKFPVLHDIGNPKAIKCVPQAEVEWKFYDAQPCSDKAYKIEELFYSYATHHDK FTDGVCLFWNCNVDRYPANAIVCRFDTRVLSNLNLPGCDGGSLYVNKHAFHTPAFDKSAF TNLKQLPFFYYSDSPCESHGKQVVSDIDYVPLKSATCITRCNLGGAVCRHHANEYRQYLD AYNMMISAGFSLWIYKQFDTYNLWNTFTRLQSLENVAYNVVNKGHFDGHAGEAPVSIINN AVYTKVDGIDVEIFENKTTLPVNVAFELWAKRNIKPVPEIKILNNLGVDIAANTVIWDYK

REAPAHVSTIGVCTMTDIAKKPTESACSSLTVLFDGRVEGOVDLFRNARNGVLITEGSVK GLTPSKGPAOASVNGVTLIGESVKTOFNYFKKVDGIIQQLPETYFTOSRDLEDFKPRSOM ETDFLELAMDEFIQRYKLEGYAFEHIVYGDFSHGQLGGLHLMIGLAKRSODSPLKLEDFI PMDSTVKNYFITDAQTGSSKCVCSVIDLLLDDFVEIIKSQDLSVISKVVKVTIDYAEISF MLWCKDGHVETFYPKLQASRAWQPGVAMPNLYKMQRMLLEKCDLONYGENAVIPKGIMMN VAKYTOLCOYLNTLTLAVPYNMRVIHFGAGSDKGVAPGTAVI,ROWI,PTGTI,I,VDSDI,NDF VSDAYSTLIGDCATVHTANKWDLIISDMYDPRTKHVTKENDSKEGFFTYLCGFIKOKLAL GGSIAVKITEHSWNADLYKLMGHFSWWTAFVTNVNASSSEAFLIGANYLGKPKEOIDGYT MHANYIFWRNTNPIOLSSYSLFDMSKFPLKLRGTAVMSLKENQINDMIYSLLEKGRLIIR ENNRVVVSSDILVNN-TNMFIFLLFLTLTSGSDLDRCTTFDDVQAPNYTQHTSSMRGVYY PDEIFRSDTLYLTQDLFLPFYSNVTGFHTINHTFGNPVIPFKDGIYFAATEKSNVVRGWV FGSTMNNKSOSVIIINNSTNVVIRACNFELCDNPFFAVSKPMGTOTHTMIFDNAFNCTFE YISDAFSLDVSEKSGNFKHLREFVFKNKDGFLYVYKGYOPIDVVRDLPSGFNTLKPIFKL PLGINITNFRAILTAFSPAODIWGTSAAAYFVGYLKPTTFMIKYDENGTITDAVDCSONP LAELKCSVKSFEIDKGIYOTSNFRVVPSGDVVRFPNITNLCPFGEVFNATKFPSVYAWER KKI SNCVADY SVLYNSTFF STFKCYGVSATKLNDLCF SNVYADSFVVKGDDVROI APGOT GVIADYNYKLPDDFMGCVLAWNTRNIDATSTGNYNYKYRYLRHGKLRPFERDISNVPFSP DGKPCTPPALNCYWPLNDYGFYTTTGIGYOPYRVVVLSFELLNAPATVCGPKLSTDLIKN QCVNFNFNGLTGTGVLTPSSKRFQPFQQFGRDVSDFTDSVRDPKTSEILDISPCAFGGVS VITPGTNASSEVAVLYQDVNCTDVSTAIHADQLTPAWRIYSTGNNVFQTQAGCLIGAEHV DTSYECDIPIGAGICASYHTVSLLRSTSOKSIVAYTMSLGADSSIAYSNNTIAIPTNFST SITTEVMPVSMAKTSVDCNMYICGDSTECANLLLQYGSFCTQLNRALSGIAAEQDRNTRE VFAOVKOMYKTPTLKYFGGFNFSQILPDPLKPTKRSFIEDLLFNKVTLADAGFMKQYGEC LGDINARDLICAOKFNGLTVLPPLLTDDMIAAYTAALVSGTATAGWTFGAGAALQIPFAM QMAYRFNGIGVTQNVLYENQKOIANQFNKAISOIOESLTTTSTALGKLODVVNONAOALN TLVKQLSSNFGAISSVLNDILSRLDKVEAEVOIDRLITGRLOSLOTYVTOOLIRAAEIRA SANLAATKMSECVLGQSKRVDFCGKGYHLMSFPQAAPHGVVFLHVTYVPSQERNFTTAPA ICHEGKAYFPREGVFVFNGTSWFITQRNFFSPQIITTDNTFVSGNCDVVIGIINNTVYDP LOPELDSFKEELDKYFKNHTSPDVDLGDISGINASVVNIQKEIDRLNEVAKNLNESLIDL QELGKYEQYIKWPWYVWLGFIAGLIAIVMVTILLCCMTSCCSCLKGACSCGSCCKFDEDD SEPVLKGVKLHYT-TNLWICL-DFLLLDQLLHSQ-KLTMLLLQVLFMLQQRYRYKPHSLS DGLLLALHFLLFFRALPK-LRSIKDGS-PFIRASSSFAIYCCYLLPSIHIFCLSLOVWRR NFCTSMP-YIFYNASTHVELL-DVGFVGSANPRTHYFMMPTTLFAGTHITMTTVYHITVS QIQLSLLKVTAFQHQNSKKTTKLVVILRIGTOVLKTMSLYMAISPKFTTSLSLHKLLOTL VLKMLHSSSLTSLLKTHRMCKYTQSTALOELLIQQWIOFMMSRRRLLACLCKHKKVSTNL CTHSFRKKQVR--LIAYFFFLLSWYSC-SH-PSLLRFDCVRTAAILLT-V--NORFTSTR VLKI-TLLKEFLIFWSKRTNYYYYSVWNFNIAYHGRORYYYR-GA-TTPGTMEPSNRFPI PSLDYVTTICLF-SEQVFVHNKACFPLALVASNTCLFCACCCLQN-LGDWRDCDCNGLYC RLDVA-LLRCFLQAVCSYPLNVVIQPRNKHSSQCASPGDNCDQTAHGK-TCHWCCDHSWS LANGRTLPRAL-H-GPAKRDHCGYITNAFLLOIRSVAACRH-FRFCCIOPLPYWKL-IKY RPRR-OROYCFASTVSDNRCFILLTSRLQ-ORY-LSL-GLSGLLFGILTL--VO--DNYL SL-LRRIIRS-MMKNLWS-IIHKTNMKIILFLTLIVFTSCELYHYOECVRGTTVLLKEPC PSGTYEGNSPFHPLADNKFALTCTSTHFAFACADGTRHTYOLRARSVSPKLFIROEEVOO ELYSPLFLIVAALVFLILCFTIKRKTE-MSSL-LTSICAF-PFCYSLF--CLLYFGFHSK SRI-KNLVPKSKRT-NFSLF-LVFLYAVAYAL-YSAVHLINLMCLKILVRYNTRGNTYST AWLCALGKVLPFHRWHTMVQTCTPNVTINCODPAGGALIARCWYLHEGHOTAAFRDVLVV LNKRTN-NV--WTPIKPT-CPPHYIWWTHRFN-O-PEWRTOWGKAKTAPTPRFTO-YCVL VHSSHSAWOGGT-IPSRPGRSNQHQ-WSR-PNWLLPKSYPTSSWW-RONERAOPOMVLLL PRNWPRSFTSLRR-QRRHRMGCN-GSLEYTORPHWHPOS--OCCHRATTSSRNNIAKRLL RRGKQRRQSSLFSLLIT-SR-FKKFNSWQQ-GKFSCSNG-RRW-NCPRAIAAROIEPA-E QSFW-RPTTTRPNCH-EICC-GI-KASPKTYCHKTVORHSSIWETWSRTNPRKFRGPRPN QTRN-LQTLAANCTICSKCLCILWNVTHWHGSHTFGNMADLSWSH-IG-ORSTIORORHT AEOAH-RIQNIPTNRA-KGOKEKD--SSAFAAETKEAAHCDSSSCG-HG-FLQTTSKFHE WSFC-FNSGINTHDDHTROMGYVNVFAIPFTIHSLLLCRMNSRN-TAOVGLVNFNLT-OS LINV-H-GGLERATTFSSRPRGVRSRVQ-IMLGRAAYMEEP-CVKLILVVLSPCDFNSFL

PTPRTLDAVCGCINVFKRVCGVSAARLTPCGTGTSTDVVYRAFDIYNEKVAGFAKFLKTNCCRFOE $\verb|KDEEGNLLDSYFVVKRHTMSNYOHEETIYNLVKDCPAVAVHDFFKFRVDGDMVPHISRORLTKYTM|$ ADLVYALRHFDEGNCDTLKEILVTYNCCDDDYFNKKDWYDFVENPDILRVYANLGERVROSLLKTV OFCDAMRDAGIVGVLTLDNODLNGNWYDFGDFVQVAPGCGVPIVDSYYSLLMPILTLTRALAAESH MDADLAKPLIKWDLLKYDFTEERLCLFDRYFKYWDQTYHPNCINCLDDRCILHCANFNVLFSTVFP PTSFGPLVRKIFVDGVPFVVSTGYHFRELGVVHNQDVNLHSSRLSFKELLVYAADPAMHAASGNLL ${ t LDKRTTCFSVAALTNNVAFOTVKPGNFNKDFYDFAVSKGFFKEGSSVELKHFFFAQDGNAAISDYD}$ YYRYNLPTMCDIRQLLFVVEVVDKYFDCYDGGCINANQVIVNNLDKSAGFPFNKWGKARLYYDSMS YEDODALFAYTKRNVIPTITQMNLKYAISAKNRARTVAGVSICSTMTNRQFHQKLLKSIAATRGAT VVIGTSKFYGGWHNMLKTVYSDVETPHLMGWDYPKCDRAMPNMLRIMASLVLARKHNTCCNLSHRF YRLANECAOVLSEMVMCGGSLYVKPGGTSSGDATTAYANSVFNICQAVTANVNALLSTDGNKIADK YVRNLQHRLYECLYRNRDVDHEFVDEFYAYLRKHFSMMILSDDAVVCYNSNYAAOGLVASIKNFKA VLYYQNNVFMSEAKCWTETDLTKGPHEFCSQHTMLVKQGDDYVYLPYPDPSRILGAGCFVDDIVKT DGTLMIERFVSLAIDAYPLTKHPNQEYADVFHLYLQYIRKLHDELTGHMLDMYSVMLTNDNTSRYW EPEFYEAMYTPHTVLQAVGACVLCNSQTSLRCGACIRRPFLCCKCCYDHVISTSHKLVLSVNPYVC NAPGCDVTDVTQLYLGGMSYYCKSHKPPISFPLCANGQVFGLYKNTCVGSDNVTDFNAIATCDWTN AGDYILANTCTERLKLFAAETLKATEETFKLSYGIATVREVLSDRELHLSWEVGKPRPPLNRNYVF ${\tt TGYRVTKNSKVOIGEYTFEKGDYGDAVVYRGTTTYKLNVGDYFVLTSHTVMPLSAPTLVPQEHYVR}$ ITGLYPTLNISDEFSSNVANYQKVGMQKYSTLQGPPGTGKSHFAIGLALYYPSARIVYTACSHAAV DALCEKALKYLPIDKCSRIIPARARVECFDKFKVNSTLEQYVFCTVNALPETTADIVVFDEISMAT NYDLSVVNARLRAKHYVYIGDPAQLPAPRTLLTKGTLEPEYFNSVCRLMKTIGPDMFLGTCRRCPA $\={\tt EIVDTVSALVYDNKLKAHKDKSAQCFKMFYKGVITHDVSSAINRPQIGVVREFLTRNPAWRKAVFI}$ SPYNSQNAVASKILGLPTQTVDSSQGSEYDYVIFTQTTETAHSCNVNRFNVAITRAKIGILCIMSD RDLYDKLQFTSLEIPRRNVATLQAENVTGLFKDCSKIITGLHPTQAPTHLSVDIKFKTEGLCVDIP GIPKDMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHLIPLMYKGLPWNVVRIKIVOMLSDTLK GLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWNHSVGFDYVYNPFM IDVOQWGFTGNLQSNHDOHCQVHGNAHVASCDAIMTRCLAVHECFVKRVDWSVEYPIIGDELRVNS ACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEWKFYDAQPCSDKAYKIEELFYSYATH + DKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSNLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPLKSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAG FSLWIYKQFDTYNLWNTFTRLQSLENVAYNVVNKGHFDGHAGEAPVSIINNAVYTKVDGIDVEIFE NKTTLPVNVAFELWAKRNIKPVPEIKILNNLGVDIAANTVIWDYKREAPAHVSTIGVCTMTDIAKK ${\tt PTESACSSLTVLFDGRVEGOVDLFRNARNGVLITEGSVKGLTPSKGPAQASVNGVTLIGESVKTQF}$ NYFKKVDGIIQQLPETYFTQSRDLEDFKPRSQMETDFLELAMDEFIQRYKLEGYAFEHIVYGDFSH GQLGGLHLMIGLAKRSQDSPLKLEDFIPMDSTVKNYFITDAQTGSSKCVCSVIDLLLDDFVEIIKS QDLSVISKVVKVTIDYAEISFMLWCKDGHVETFYPKLQASRAWQPGVAMPNLYKMQRMLLEKCDLO NYGENAVIPKGIMMNVAKYTQLCQYLNTLTLAVPYNMRVIHFGAGSDKGVAPGTAVLRQWLPTGTL ${\tt LVDSDLNDFVSDAYSTLIGDCATVHTANKWDLIISDMYDPRTKHVTKENDSKEGFFTYLCGFIKOK}$ LALGGSIAVKITEHSWNADLYKLMGHFSWWTAFVTNVNASSSEAFLIGANYLGKPKEQIDGYTMHA ${\tt NYIFWRNTNPIQLSSYSLFDMSKFPLKLRGTAVMSLKENQINDMIYSLLEKGRLIIRENNRVVVSS}$ ${\tt DILVNN\underline{*}TNMFIFLLFLTLTSGSDLDRCTTFDDVQAPNYTQHTSSMRGVYYPDEIFRSDTLYLTQD}$ LFLPFYSNVTGFHTINHTFGNPVIPFKDGIYFAATEKSNVVRGWVFGSTMNNKSQSVIIINNSTNV VIRACNFELCDNPFFAVSKPMGTQTHTMIFDNAFNCTFEYISDAFSLDVSEKSGNFKHLREFVFKN KDGFLYVYKGYQPIDVVRDLPSGFNTLKPIFKLPLGINITNFRAILTAFSPAQDIWGTSAAAYFVG ${\tt YLKPTTFMLKYDENGTITDAVDCSQNPLAELKCSVKSFEIDKGIYQTSNFRVVPSGDVVRFPNITN}$ LCPFGEVFNATKFPSVYAWERKKISNCVADYSVLYNSTFFSTFKCYGVSATKLNDLCFSNVYADSF VVKGDDVRQIAPGQTGVIADYNYKLPDDFMGCVLAWNTRNIDATSTGNYNYKYRYLRHGKLRPFER DISNVPFSPDGKPCTPPALNCYWPLNDYGFYTTTGIGYQPYRVVVLSFELLNAPATVCGPKLSTDL ${\tt IKNQCVNFNFNGLTGTGVLTPSSKRFQPFQQFGRDVSDFTDSVRDPKTSEILDISPCAFGGVSVIT}$ PGTNASSEVAVLYQDVNCTDVSTAIHADQLTPAWRIYSTGNNVFQTQAGCLIGAEHVDTSYECDIP IGAGICASYHTVSLLRSTSQKSIVAYTMSLGADSSIAYSNNTIAIPTNFSISITTEVMPVSMAKTS $ext{VDCNMYICGDSTECANLLLQ}{ imes} ext{GSFCTQLNRALSGIAAEQDRNTREVFAQVKQMYKTPTLKYFGGFN}$ ${ t FSQILPDPLKPTKRSFIEDLLFNKVTLADAGFMKQYGECLGDINARDLICAQKFNGLTVLPPLLTD}$

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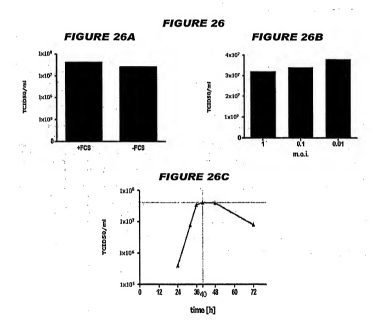
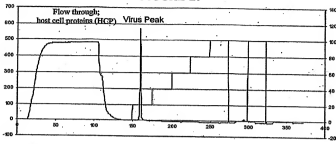


FIGURE 27



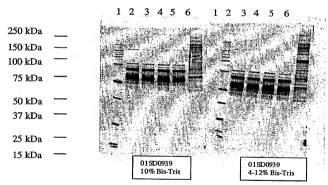
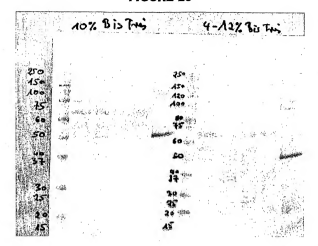
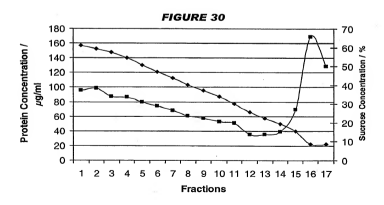


FIGURE 29





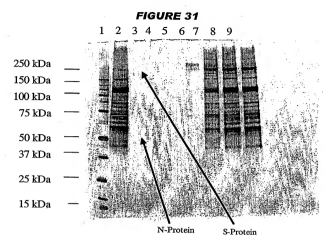


FIGURE 32

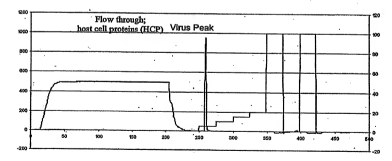


FIGURE 33

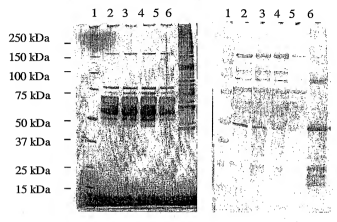
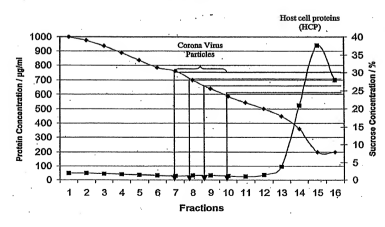
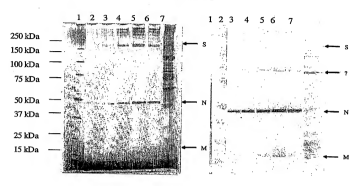
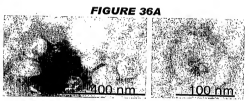


FIGURE 34







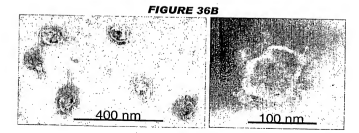
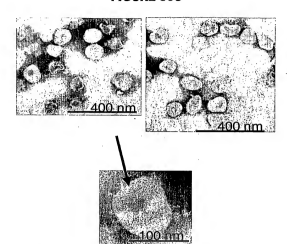


FIGURE 36C



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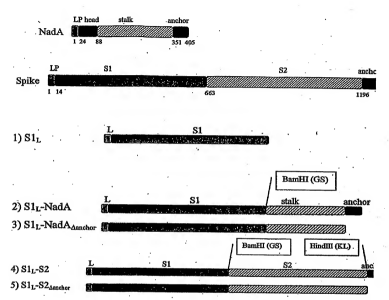


FIGURE 38

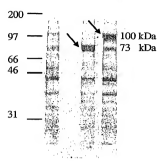
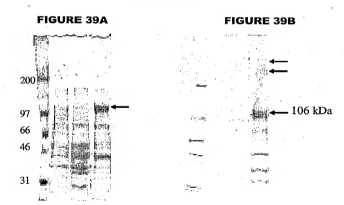
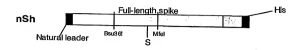
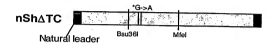
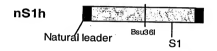


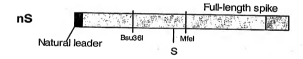
FIGURE 39

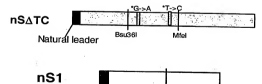














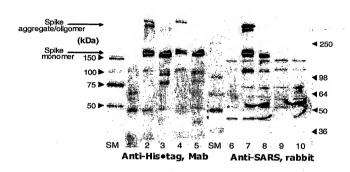
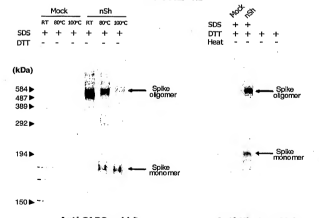
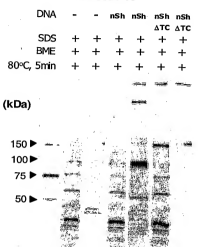


FIGURE 42

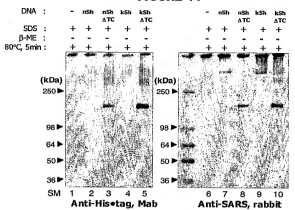


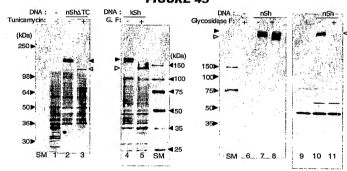
Anti-SARS, rabbit

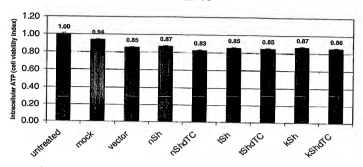
Anti-His•tag, Mab



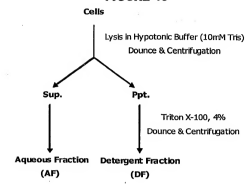
SM



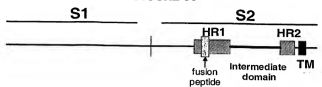




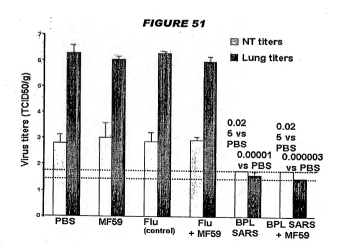
	_	Moc	<u>k_</u>	_				nSh	_				_				Mod	k				nSh		
Fraction SDS DTT Heat	9 +	10 + +	11 +	7 + + -	+ +	9 + +	10 + - -	11 +	12 + +	13 + + -	14 + -	15 + + -	16 + +	Fraction SDS DTT 80°C, 5min	+	11 + +	12 + +	13 + + +	14 + + +	10 + + +	11 + +	12 + + +	13 + +	14 + +
584 ► 487 ►									4					(kDa) 584▶ 487▶ 389▶										
							-	A.E.	,e.					194▶										
150►:.						•								150▶					ř.	entil 1	17.93	1.000		



<u>L</u> P	S1	NadA-sta	lk anchor
<u>\$</u>			
1 29/14		662 /88	405
LP	S1	NadA-sta	alk
1 29/14		662 /88	350

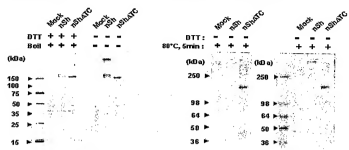


- a) Leader(NadA)-HR1-GGGGGG-HR2-GGGGSG-stalk(NadA)-anchor(NadA)
- b) Leader(NadA)-HR1-GGGGGG-HR2-GGGGSG-stalk(NadA)
- c) Leader(NadA)-HR1-intermediate-domain-HR2-GGGGSG-stalk(NadA)-anchor(NadA)
- d) Leader(NadA)-HR1-intermediate-domain-HR2- GGGGSG-stalk(NadA)
- e) HR1-intermediate-domain-HR2- GGGGSG-stalk(NadA) HHHHHH
- f) Leader(NadA)-HR1-intermediate-domain-HR2-anchor(NadA)
- g) Leader(NadA)-HR1-intermediate-domain-HR2



A. 293 cell lysates

B. COS7 cell culture supernatants



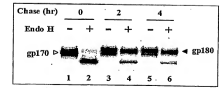
Anti-His • tag, mAb

Anti-His • tag, mAb Ant

Anti-SARS, rabbit

4-20% TG SDS gel **FIGURE 53** 4-20% TG SDS gel

Mock		nSh	١		nSh/	MC		
RT 90°C 100°C	RTE	0°C	100*0	RT	80°C	100*	С	
		4	· · · · · · · · · · · · · · · · · · ·	garan Salah Palah	1 m		4	669
	Tuber of	Car	*	-	4		4	440
						,	4	232
<u>.</u>					C.	ĝ. ()		
							4	158



107/193

FIGURE 54

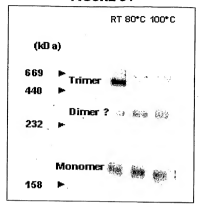
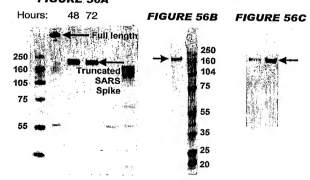
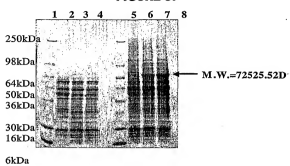
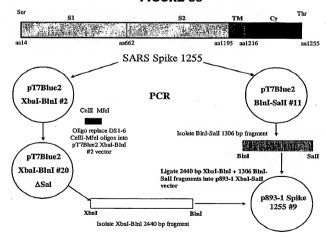


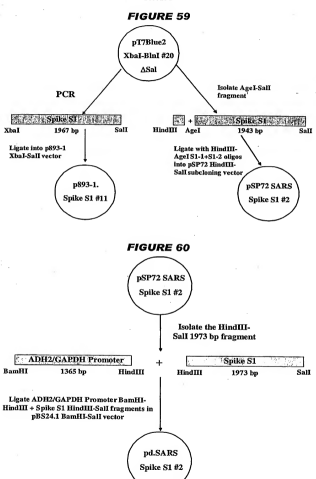
FIGURE 56

FIGURE 56A

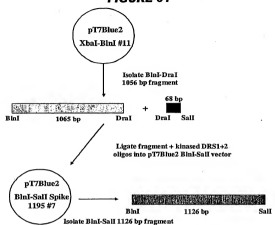


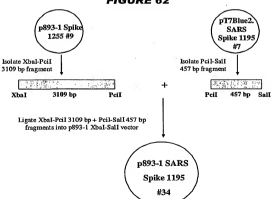


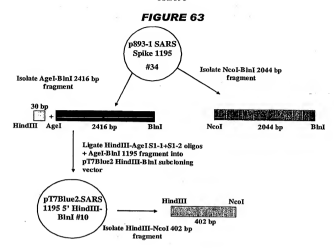


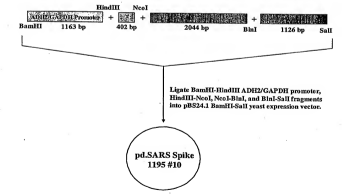












AAGO	TTAC	CAAAZ	ACAA	Y	1 M ATG	S AGT	D GAC	L CTT			C TGC			
D GAT		V GTT		A GCT				T ACT		H CAT		S TCA	S TCT	M ATG
R AGG	G GGG	V GTT	Y TAC	30 Y TAT		D GAT		I ATT				D GAC	T ACT	40 L CTT
Y TAT	L TTA	T ACT	Q CAG	D GAT			L CTT	P CCA	50 F TTT	Y TAT	S TCT	N AAT	V GTT	T ACA
G GGG	F TTT	H CAT	T ACT	60 I ATT	N AAT	H CAT	T ACG	F TTT	G GGC	N AAC		V GTC	I ATA	70 P CCT
F TTT	K AAG	D GAT	G GGT		Y TAT		A GCT	A GCC	BO T ACA	E GAG	K AAA	S TCA	N AAT	V GTT
V GTC	R CGT	G GGT	W TGG		F			T ACC			N AAC			100 Q CAG
S TCG	V GTG			I ATT				T ACT	110 N AAT	V GTT	V GTT	I ATA		A GCA
C TGT	N AAC	F TTT	E GAA	120 L TTG	C TGT			P CCT						
P	M ATG	G GGT		Q CAG			T ACT	M ATG					A GCA	F TTT
N AAT	C TGC	T ACT		150 E GAG				D GAT				L CTT		160 V GTT
S TCA	E GAA			G GGT				H CAC		R			V GTG	F TTT
K AAA	N AAT	K AAA	D GAT	180 G GGG	F TTT		Y TAT	V GTT			_		~	190 P CCT

200 V V R D L P S G F N T L K D I ATA GAT GTA GTT CGT GAT CTA CCT TCT GGT TTT AAC ACT TTG AAA 210 220 F L GINITN CCT ATT TTT AAG TTG CCT CTT GGT ATT AAC ATT ACA AAT TTT AGA 230 AFSPA 0 I D GCC ATT CTT ACA GCC TTT TCA CCT GCT CAA GAC ATT TGG GGC ACG 240 250 Y F V G Y L K P F TCA GCT GCA GCC TAT TTT GTT GGC TAT TTA AAG CCA ACT ACA TTT 260 L к у р E N G T I D ATG CTC AAG TAT GAT GAA AAT GGT ACA ATC ACA GAT GCT GTT GAT 270 280 SONP L AELK C S TGT TCT CAA AAT CCA CTT GCT GAA CTC AAA TGC TCT GTT AAG AGC 290 IDKGIYQT N TTT GAG ATT GAC AAA GGA ATT TAC CAG ACC TCT AAT TTC AGG GTT 300 310 V P S G D V V R F P N I T L GTT CCC TCA GGA GAT GTT GTG AGA TTC CCT AAT ATT ACA AAC TTG 320 P F G E V F N A T TGT CCT TTT GGA GAG GTT TTT AAT GCT ACT AAA TTC CCT TCT GTC 330 340 R K K I S N C V TAT GCA TGG GAG AGA AAA AAA ATT TCT AAT TGT GTT GCT GAT TAC 350 LYNSTFF S T TCT GTG CTC TAC AAC TCA ACA TTT TTT TCA ACC TTT AAG TGC TAT 360 370 T LNDLCFS A K v GGC GTT TCT GCC ACT AAG TTG AAT GAT CTT TGC TTC TCC AAT GTC 380 SFVVKG D D TAT GCA GAT TCT TTT GTA GTC AAG GGA GAT GAT GTA AGA CAA ATA . 390 400 T G V I A D Y N Y GCG CCA GGG CAA ACT GGT GTT ATT GCT GAT TAT AAT TAT AAA TTG

410 PDDFMGCVLA WNTR CCA GAT GAT TTC ATG GGT TGT GTC CTT GCT TGG AAT ACT AGG AAC 420 430 T S T G N Y N Y K Y R Y ATT GAT GCT ACT TCA ACT GGT AAT TAT AAT TAT AAA TAT AGG TAT 440 HGKLRPF R Ε D I S CTT AGA CAT GGC AAG CTT AGG CCC TTT GAG AGA GAC ATA TCT AAT DGKPCTPP P GTG CCT TTC TCC CCT GAT GGC AAA CCT TGC ACC CCA CCT GCT CTT 470 CYWPLNDYGFYTTT AAT TGT TAT TGG CCA TTA AAT GAT TAT GGT TTT TAC ACC ACT ACT 480 490 Y YRVVVLS 0 GGC ATT GGC TAC CAA CCT TAC AGA GTT GTA GTA CTT TCT TTT GAA 500 PATVCGP CTT TTA AAT GCA CCG GCC ACG GTT TGT GGA CCA AAA TTA TCC ACT 510 520 K N O C V N F N F N G L I GAC CTT ATT AAG AAC CAG TGT GTC AAT TTT AAT TTT AAT GGA CTC 530 G T G V L T P S S K ACT GGT ACT GGT GTG TTA ACT CCT TCT TCA AAG AGA TTT CAA CCA 540 550 RDVSDFTD G TTT CAA CAA TTT GGC CGT GAT GTT TCT GAT TTC ACT GAT TCC GTT 560 PKTSEI L D I S CGA GAT CCT AAA ACA TCT GAA ATA TTA GAC ATT TCA CCT TGC TCT 570 V S V I T P G T N A S TTT GGG GGT GTA AGT GTA ATT ACA CCT GGA ACA AAT GCT TCA TCT 590 V A V L Y Q D V N C T D V S GAA GTT GCT GTT CTA TAT CAA GAT GTT AAC TGC ACT GAT GTT TCT 600 610 IHADOLTPAWR ACA GCA ATT CAT GCA GAT CAA CTC ACA CCA GCT TGG CGC ATA TAT

G, A G I C A S Y H T OC SEQ ID NO: 9799
GGA GCT GGC ATT TGT GCT AGT TAC CAT ACA TAA TGAGTCGAC SEQ ID NO: 9800
Translated Mol. Weight = 72525.52

FIGURE 66

1 10 D R C AAGCTTACAAAACAAA ATG AGT GAC CTT GAC CGG TGC ACC ACT TTT 20 P Y N Н GAT GAT GTT CAA GCT CCT AAT TAC ACT CAA CAT ACT TCA TCT ATG 30 Y P E I F R AGG GGG GTT TAC TAT CCT GAT GAA ATT TTT AGA TCA GAC ACT CTT L L F TAT TTA ACT CAG GAT TTA TTT CTT CCA TTT TAT TCT AAT GTT ACA 60 I N т G N GGG TTT CAT ACT ATT AAT CAT ACG TTT GGC AAC CCT GTC ATA CCT Y F TTT AAG GAT GGT ATT TAT TTT GCT GCC ACA GAG AAA TCA AAT GTT 100 v s M N GTC CGT GGT TGG GTT TTT GGT TCT ACC ATG AAC AAC AAG TCA CAG 110 I I N N S N TCG GTG ATT ATT AAC AAT TCT ACT AAT GTT GTT ATA CGA GCA 130 N P F F TGT AAC TTT GAA TTG TGT GAC AAC CCT TTC TTT GCT GTT TCT AAA 140 0 T н М т CCC ATG GGT ACA CAG ACA CAT ACT ATG ATA TTC GAT AAT GCA TTT 150 E Y I S D A s AAT TGC ACT TTC GAG TAC ATA TCT GAT GCC TTT TCG CTT GAT GTT

170 F K S G N F K H L R TCA GAA AAG TCA GGT AAT TTT AAA CAC TTA CGA GAG TTT GTG TTT 180 LYVYKGY D G AAA AAT AAA GAT GGG TTT CTC TAT GTT TAT AAG GGC TAT CAA CCT 200 V V R D L P S G FNTL ATA GAT GTA GTT CGT GAT CTA CCT TCT GGT TTT AAC ACT TTG AAA 210 220 LGINITNF F ĸ L P CCT ATT TTT AAG TTG CCT CTT GGT ATT AAC ATT ACA AAT TTT AGA 230 AFSP I W G T A O D GCC ATT CTT ACA GCC TTT TCA CCT GCT CAA GAC ATT TGG GGC ACG 240 250 YFVGYLKPT Α TCA GCT GCA GCC TAT TTT GTT GGC TAT TTA AAG CCA ACT ACA TTT 260 LKYDENGT I TDAVD ATG CTC AAG TAT GAT GAA AAT GGT ACA ATC ACA GAT GCT GTT GAT 270 280 PLAELKCSV 0 N TGT TCT CAA AAT CCA CTT GCT GAA CTC AAA TGC TCT GTT AAG AGC 290 E. I D K G I Y O S N F R V T TTT GAG ATT GAC AAA GGA ATT TAC CAG ACC TCT AAT TTC AGG GTT 300 310 S G VVRFPNTTN D GTT CCC TCA GGA GAT GTT GTG AGA TTC CCT AAT ATT ACA AAC TTG 320 PFGEVFN KFPS A T TGT CCT TTT GGA GAG GTT TTT AAT GCT ACT AAA TTC CCT TCT GTC 330 340 R K K I S N C V E TAT GCA TGG GAG AGA AAA AAA ATT TCT AAT TGT GTT GCT GAT TAC 350 YNSTFF s TCT GTG CTC TAC AAC TCA ACA TTT TTT TCA ACC TTT AAG TGC TAT 360 370 T KLNDLCFS GGC GTT TCT GCC ACT AAG TTG AAT GAT CTT TGC TTC TCC AAT GTC 380 SFVVK G D D V R TAT GCA GAT TCT TTT GTA GTC AAG GGA GAT GAT GTA AGA CAA ATA 390 400 TGVIADYNY GCG CCA GGG CAA ACT GGT GTT ATT GCT GAT TAT AAT TAT AAA TTG

410 CVL М G Α CCA GAT GAT TTC ATG GGT TGT GTC CTT GCT TGG AAT ACT AGG AAC 420 430 s T G N Y N Y ĸ ATT GAT GCT ACT TCA ACT GGT AAT TAT AAT TAT AAA TAT AGG TAT L R P F E R D CTT AGA CAT GGC AAG CTT AGG CCC TTT GAG AGA GAC ATA TCT AAT 450 460 D G ĸ P C т P P GTG CCT TTC TCC CCT GAT GGC AAA CCT TGC ACC CCA CCT GCT CTT P L N D Y G F Y AAT TGT TAT TGG CCA TTA AAT GAT TAT GGT TTT TAC ACC ACT ACT 480 490 R V V V Y L GGC ATT GGC TAC CAA CCT TAC AGA GTT GTA GTA CTT TCT TTT GAA 500 v C A G CTT TTA AAT GCA CCG GCC ACG GTT TGT GGA CCA AAA TTA TCC ACT 510 520 ĸ N v NFNFN GAC CTT ATT AAG AAC CAG TGT GTC AAT TTT AAT TTT AAT GGA CTC T P S s ACT GGT ACT GGT GTG TTA ACT CCT TCT TCA AAG AGA TTT CAA CCA 550 G V S D F R D T TTT CAA CAA TTT GGC CGT GAT GTT TCT GAT TTC ACT GAT TCC GTT 560 KTSEI L D . І s CGA GAT CCT AAA ACA TCT GAA ATA TTA GAC ATT TCA CCT TGC TCT 570 580 т P G T N TTT GGG GGT GTA-AGT GTA ATT ACA CCT GGA ACA AAT GCT TCA TCT 590 v L Y 0 D v N С GAA GTT GCT GTT CTA TAT CAA GAT GTT AAC TGC ACT GAT GTT TCT 600 610 Α D L TPAWR ACA GCA ATT CAT GCA GAT CAA CTC ACA CCA GCT TGG CGC ATA TAT 620 N N v F т 0 TCT ACT GGA AAC AAT GTA TTC CAG ACT CAA GCA GGC TGT CTT ATA 630 H V D T s Y Е C D Ι

GGA GCT GAG CAT GTC GAC ACT TCT TAT GAG TGC GAC ATT CCT ATT 650 ICASYHT GGA GCT GGC ATT TGT GCT AGT TAC CAT ACA GTT TCT TTA TTA CGT 660 K S I V A Y T M S L AGT ACT AGC CAA AAA TCT ATT GTG GCT TAT ACT ATG TCT TTA GGT 680 DSSIAYS I A I N N GCT GAT AGT TCA ATT GCT TAC TCT AAT AAC ACC ATT GCT ATA CCT 690 700 ISITTEVMP ACT AAC TTT TCA ATT AGC ATT ACT ACA GAA GTA ATG CCT GTT TCT 710 K T S V D C N M Y I C G ATG GCT AAA ACC TCC GTA GAT TGT AAT ATG TAC ATC TGC GGA GAT 720 730 С A N L L L Q Y G S TCT ACT GAA TGT GCT AAT TTG CTT CTC CAA TAT GGT AGC TTT TGC 740 OLNRALS G I ACA CAA CTA AAT CGT GCA CTC TCA GGT ATT GCT GCT GAA CAG GAT 750 760 R EVFAQVKQMY CGC AAC ACA CGT GAA GTG TTC GCT CAA GTC AAA CAA ATG TAC AAA 770 T P T L K Y F G G F N F S O ACC CCA ACT TTG AAA TAT TTT GGT GGT TTT AAT TTT TCA CAA ATA 780 790 P D L K P T K R S F T TTA CCT GAC CCT CTA AAG CCA ACT AAG AGG TCT TTT ATT GAG GAC 800 F N K V T L Α D A TTG CTC TTT AAT AAG GTG ACA CTC GCT GAT GCT GGC TTC ATG AAG 810 820 L GDINARD E C CAA TAT GGC GAA TGC CTA GGT GAT ATT AAT GCT AGG GAC CTC ATT 830 K F N G L T v L TGT GCG CAG AAG TTC AAT GGA CTT ACA GTG TTG CCA CCT CTG CTC 840 850 M I Y ACT GAT GAT ATG ATT GCT GCC TAC ACT GCT GCT CTA GTT AGT GGT 860 G WTFG A ACT GCC ACT GCT GGA TGG ACA TTT GGT GCT GGC GCT GCT CTT CAA 870 880

M А Y R ATA CCT TTT GCT ATG CAA ATG GCA TAT AGG TTC AAT GGC ATT GGA 890 V L Y E N GTT ACC CAA AAT GTT CTC TAT GAG AAC CAA AAA CAA ATC GCC AAC 900 K I S O I O E S T. CAA TTT AAC AAG GCG ATT AGT CAA ATT CAA GAA TCA CTT ACA ACA 920 L G K L D ACA TCA ACT GCA TTG GGC AAG CTG CAA GAC GTT GTT AAC CAG AAT 930 L N T VKQLSS L GCT CAA GCA TTA AAC ACA CTT GTT AAA CAA CTT AGC TCT AAT TTT 950 S v L N D I GGT GCA ATT TCA AGT GTG CTA AAT GAT ATC CTT TCG CGA CTT GAT 960 E 7.7 IDRLTT AAA GTC GAG GCG GAG GTA CAA ATT GAC AGG TTA ATT ACA GGC AGA 980 т Y V T CTT CAA AGC CTT CAA ACC TAT GTA ACA CAA CAA CTA ATC AGG GCT 990 N L A A T K Α А GCT GAA ATC AGG GCT TCT GCT AAT CTT GCT GCT ACT AAA ATG TCT 1010 G S ĸ R V D GAG TGT GTT CTT GGA CAA TCA AAA AGA GTT GAC TTT TGT GGA AAG 1020 M A GGC TAC CAC CTT ATG TCC TTC CCA CAA GCA GCC CCG CAT GGT GTT 1040 L H V T Y v Ρ S GTC TTC CTA CAT GTC ACG TAT GTG CCA TCC CAG GAG AGG AAC TTC 1050 Α I C HEGKAYF ACC ACA GCG CCA GCA ATT TGT CAT GAA GGC AAA GCA TAC TTC CCT 1070 F V F N G CGT GAA GGT GTT TTT GTG TTT AAT GGC ACT TCT TGG TTT ATT ACA 1080 S F P QIITTDN CAG AGG AAC TTC TTT TCT CCA CAA ATA ATT ACT ACA GAC AAT ACA 1100 SGNCDVVI TTT GTC TCA GGA AAT TGT GAT GTC GTT ATT GGC ATC ATT AAC AAC

1140 1150 F G D I S G I N A S V V N I Q

TTT GGC GAC ATT TCA GGC ATT AAC GCT TCT GTC GTC AAC ATT CAA

v

A K N

AAA GAA ATT GAC CGC CTC AAT GAG GTC GCT AAA AAT TTA AAT GAA 1170

S L I D L Q E L G K Y E Q Y I TCA CTC ATT GAC CTT CAA GAA TTG GGA AAA TAT GAG CAA TAT ATT

1183

K W P OC SEQ ID NO: 9801

AAA TGG CCT TAA TGAGTCGAC SEQ ID NO: 9802

Translated Mol. Weight = 131315.20

EIDRLNE

FIGURE 67

FIGURE 67A

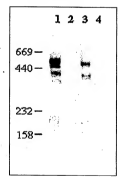
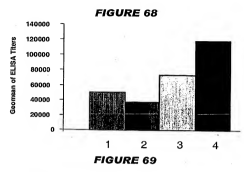


FIGURE 67B

	Virian
	RT 80°C 100°C
(kD	a)
669 440	Trimer
232	Dimer? as 200 (d.
158	Monomer) & (a)



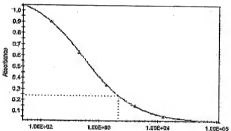


FIGURE 70

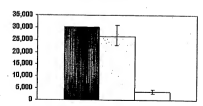
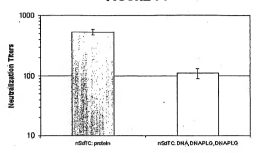
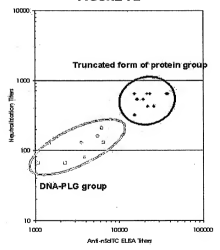


FIGURE 71







1 2

FIGURE 73

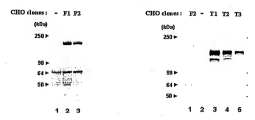
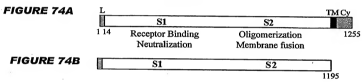


FIGURE 74



F	igui	RE 7	75A				FIGURE 75B				
	ź	ock	₁₁ Sh	4	BHATC	5		Mockinst	1 ISHATC		
PNGase F	: -	 + -	+	_	+		DTT:		_		
(kDa)							80°C, 5min:	+ +	+		
250 —		· 					(kDa)				
150 —	,	i to	١			•	250 —				
100 —		•		. '	-	•			**** ■		
75 —											
50 —					٠	ř	98 —				
35—							64 —				
25 -							50 —				
	1 2	3	4	5	6		36—				

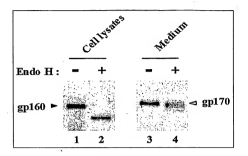


FIGURE 77

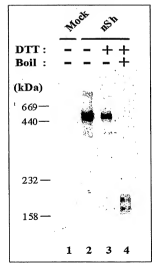


FIGURE 78

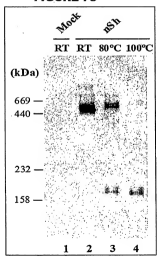


FIGURE 79

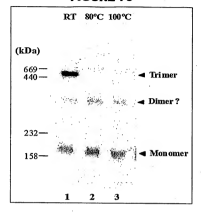


FIGURE 80

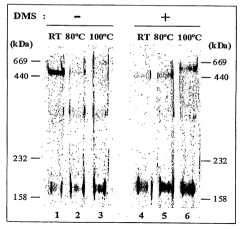


FIGURE 81

RT 80°C 100°C

FIGURE 82

RT 80°C 100°C

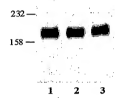
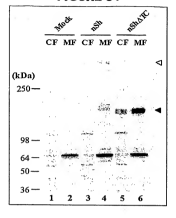
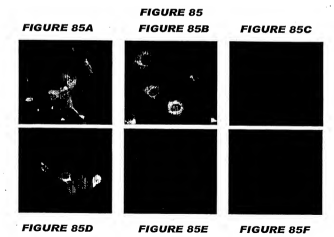




FIGURE 84







P28 (19.7)(Da)

FIGURE 87

P65 PET Tot sol

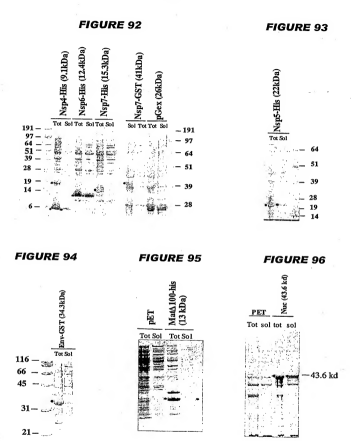
FIGURE 88

210 — Tot 1111 — 71 — 55 — 41 — 41

FIGURE 89



FIGURE 90







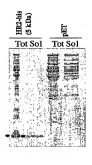


FIGURE 99

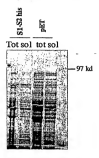


FIGURE 100



FIGURE 101



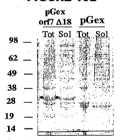


FIGURE 103



FIGURE 104



FIGURE 105

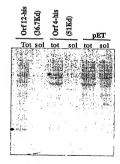


FIGURE 106

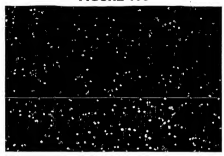


FIGURE 107

FIGURE 107A

FIGURE 107B



FIGURE 108

FIGURE 108A



FIGURE 108B



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FIGURE 109

S1 S1-S2

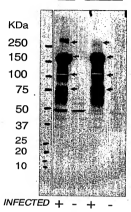
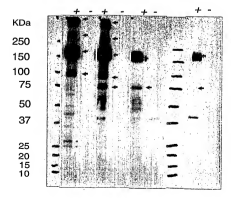
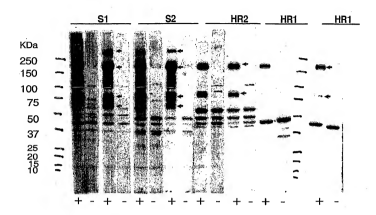
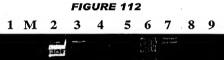


FIGURE 110 S1 S1-S2 HR2

<u>HR1</u>









5'3' Frame 1

PKDMTYVDSSL-WVS-ITKSMVTLICLSPAKKLFVTFVRGLALM-RAVMQLEMLWVLTYL SS-DFLQVLT--LYRLVMLTLKITQNSPELMHKPPPVSSLNILYHSCIKACPGM-CVLR-YKCSVIH-KDCQTESCSSFGRMALSLHQ-STLSRLDLKERVVCVTNVQLAFLLHQILMPA GIILWVLTMSITHL-LMFSSGGFTGNLSE-P-PTLPGTWKCTCGLVVMI.

5'3' Frame 2

QRT-PT-THLYDGFQNELPSQWLP-YVYHPRRSYSSRSCVDWL-CRGLSCN-RCCGY-PT SPARIFYRC-LSSCTDWLC-H-K-HKIHQS-CTNLHQ-AV-TSYTTHV-RLALECSAY-D STNAQ-YTERIVRQSRVRPLGAWL-AYINEVLCQDWT-KNVLSV-QTCNLLFYFIRYLCL LESFCGF-LCL-PIYD-CSAVGALRVTFQSNHDOHCOVHGNAHVG-L-C

5'3' Frame 3

KGHDLRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLP LQLGFSTGVNLVAVPTGYVDTENNTKFTRVNAQTSTSEQFKHLIPLMYKGLPWNVVRIKI VQMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYAC WNHSVGFDYVYNPFMIDVQQWGLYG-PFRVTMTNIARYMEMHMWASCDA

3'5' Frame 1

-HHN-PTCAFPCTWQCWSWLL-KVTRKAPTAEHQS-MGYRHSQNPQNDSSRHKYLMK-KS KLHVCHTDNTFFQVQS-QSTSLM-AQSHAPKGRTRLCLTTLSVYH-AFVLS-YALHSRAS LYT-VV-DV-TAHWWRFVH-LW-ILCYFQCQHNQSVQLLS-HL-KILAGEVG-YPQHL-L HDSPLHQSQSTHERDE-LLRG-TY-GNH-LGNSF-NPS-R-VYVGHVLW

3'5' Frame 2

SITTSPHVHFHVPGNVGHGYSERLPVKPPLLNINHKWVIDIVKTHRMIPAGISI--SRKA SCTFVTQTTRSFRSNLDKVLH-CKLKAMRPKDEHDSV-QSFQCITEHLYYLNTHYIPGQA FIHEWYKMFKLLTGGGLCINSGEFCVIFSVNITSRYSY-VNTCRKS-LER-VSTHSISSC MTALYIKANPRTNVTNSFFAGDKHIRVTIDLVIHFETHHRDEST-VMSF

3'5' Frame 3

ASQLAHMCISMYLAMLVMVTLKGYP-SPHC-TSIINGL-T-SKPTE-FQQA-VSDEVEKQ VARLSHRQHVLSGPILTKYFIDVSSKPCAQRTNTTLSDNPFSVSLSICTILIRTTFQGKP LYMSGIRCLNCSLVEVCALTLVNFVLFSVST-PVGTATKLTPVENPSWRGRLVPTASLVA -QPSTSKPIHART-RIASSRVINILG-PLTW-FILKPIIEMSLRRSCPL

5'3' Frame 1

YRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQLG FSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHI.T

5'3' Frame 2

TVDSSL-WVSK-ITKSMVTLICLSPAKKLFVTFVRGLALM-RAVMQLEMLWVLTYLSS-D FLOVLT--LYRLVMLTLKITONSPELMONLHOVTSLNILY

5'3' Frame 3

P-THLYDGFQNELPSQWLP-YVYHPRRSYSSRSCVDWL-CRGLSCN-RCCGY-PTSPARI FYRC-LSSCTDWLC-H-K-HRIHOS-CKTSTR-PV-TSYT

3'5' Frame 1

GIRCLNWSPGGGFALTLVNSVLFSVST-PVGTATKLTPVENPSWRGRLVPTASLVA-QPS.
TSKPIHART-RIASSRVINILG-PLTW-FILKPIIEMSLR

3'5' Frame 2

V-DV-TGHLVEVLH-LW-ILCYFQCQHNQSVQLLS-HL-KILAGEVG-YPQHL-LHDSPL HQSQSTHERDE-LLRG--TY-GNH-LGNSF-NPS-R-VYG

3'5' Frame 3

YKMFKLVTWWRFCINSGEFCVIFSVNITSRYSY-VNTCRKS-LER-VSTHSISSCMTALY IKANPRTNVTNSFFAGDKHIRVTIDLVIHFETHHRDESTV

BNSDOCID: <WO____2004092380A2_1_>

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		·····					s	ection 151
(5851)	5851	586	j	5	370			588
(5675)	LTHY	LSVIKA	LIRAR	HYVY	ICDPA	JLP.	APRV	LLEKGT
(3247)	TITALE	SLSFING	CINYC	YVVY	VGDFA	OLP.	APP	五五五五二十二五五五五五五五五五五五五五五五五五五五五五五五五五五五五五五五五
(5762)	LTMY	LEVINS	RVSAF	HYVY	IGDPA	OLP.	APPU	TINECT
(1)								
(5851)	LTNY	ELSVINA	RI AI	CHYVY	IGDPA	OLP	APRV	T.T.NKGT
								Section 152
(5890)			00		5910			592
(5714)	EPRY	ENTVIKE	MCCLC	PDIF	LGTCY	ROP	KETV	DTVSAL
(5285)	SPKD	YNVVTNL	IVCVE	POTE	TAKCY	R (P	K P T W	Duright
(5801)	EPRY	FNSVTKL	ACCT.	POTE	TOTICY	о с о	E E T T	
(1)	ALAR MARKET	A CONTRACTOR OF THE PARTY OF TH	Carried St. of S	NAT PARTIES TANK				
	EPKY	FNSVTKL	MCCLG	PDIF	LGTCY	RCP	KETV	DTVSAL
								ection 153
(5929)			940		5950			596
(5753)	PENK	lkannes.	SILCE	KWYY	KG	VIT	HESS	SAVNMO
(5324)	TOGK	FIRMNER	RECI	KNIV	NEGNS	DVG	HE SIC	SAYNTH
(5840)	YOUR	CKARUDN.	SNIG	KVYY	xā	ייינים	HESS	SALUUM
` (1)		- Commercial days - o	A-0.55 CH 7850-2-10	200				Mark British
	YDNK	LKAKND	SSLCE	KVYY	KG	Trop	HESS	SAVNMO
								Section 154
(5968)			5980		5990			600
(5789)	IXLI	NKELKAN	PLVHE	AVET	SPYNS	ONE	AAKR	VLCLOT
(5363)	LEFT	KDEVCRN	KOVERE	AIFI	SEYNA	MNO	RAYR	MIGINV
(5876)	THEI	SKELKAN	PSWEI	IAVET	SPYNS	S N Y	VAKE	TELETAT
` (1)	The state of the s							
	IHLI	KFLKAN	PW	AVFI	SPYNS	ONE	AKR	VLGLQT
								Section 155
(6007)	6007		6020	Ň.	603	n		604
		AQGSEYD						
		SOGSEYD						
		AQGSEYD	FATA	SUTAE	TAHUV	M A III	P.F. DI V	ALTRAK
(1)								
(6007)	TVDS	AQGSEYD	YVIY	SQTAE	TAHSV	NVN		
				-				Section 156
(6046)				60		070		T→ 60E
(5867)	GILC	VMSNMQL	FEAL	2FTTI	TLDKV	PQA	VETR	VOCSTN
		VMRQRUE						
		VMSSMQD						
` (1 ³								
(6046	GILC	VMSNMQL	FESL	NETTI	TLDKI		P	LOCSTN

FIGURE 115 (contd.)

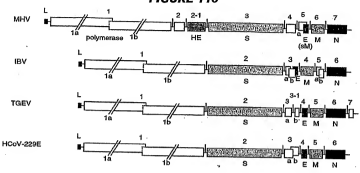
					Section	on 157
(6085) (5085	6090	6100	6110	·	6123
5906)	EKDC	SKSYSGY!	IBAHAPSEL.	AVDDKYKA	TGDLAVE	LGIG
			IPAYAVITK			
∮ (5990) - (1)		SRSYVGYI	ipahapseli 	awookykv	GCDLAVC	LNVA
(6085)	FKDC	SKSYSGYI	HPAHAPSFL	AVDDKYKV	GDLAVO	
(6124)	6494	6130	6140	AX		9
(6124)				615		6162
2242)	U-5.4	VIII SPIL	ion from			VKH
			illi ficmsv			
			LMDESTOR			
(1)	к <u>с</u>	HDLRE	Markering.	DAMASTEND	T. R. E.	TEF
(6124)	D SA	VTYSRLI	SLMGFKLDV	TLDGYCNL	FITRDE F	LIKRV on 159
(6163)	6163	6170	6180		190	6201
59831	AUT	COMERGN	REFERENCE TO	AF BLOID	SWITTER.	VEAT
(5551)	BOLV	STIVEAT	TOCGINIST	HE Provin	STUADE	WTPE
(6067)	ALTU	ge dae ga	aatrošto:	is a situation	STATE	VEAT
(38)	rATT	70000	: TRDAVIT	ELeboic v	STEVAL	VPT
(6163)	RAWV	GPDVEGA	HATROSIGT	nlplolge	STGIDET ——Secti	VEPI
(6202)		,6210	,622		6230	6240
			KKAVAKAEE			
			EPVNSKAFP			
(6106)	MEA	ERDGYV	KKA AARA PP	GENHANDED	P MSRG	rne)
			TRUBACTST			
(6202)	GLVD	TRDGY F	KKVNAKAPP	GEQFKHLI	PLMSRG	
(6241)	6241	625	0 6	260		627
		HANNEAG	HLIDISTC	LADIAAL	FELTCLI	T.FA
			NICHVSOCT			
(6145)		ESTERNIST STATE	HUADLANS	'1 TUAAS	ELTCL	a coat
(6145)	VEIF	evon iš i Živomuso	HLADLANS V	VENTUAAS VENTUAHO	ise lyck Te lysk	r i cai
(116)	VE.IF	OBLINE VI	PLECLSERY	WENTHE	TE L'ESM	() - V
(116)	VE.IF	OBLINE VI	HLADLANS PLRCLSERV HL DLSDCV	WENTHE	TE L'ESM	RYFV
(116) (6241) (6280)	VEIF VEIF VRPF	OBJENOVIJ DAJMOVIS 8	PLEGLSPER HL DLSDCV 290	VVLVTWAHC	FELTCL Sect	RYFV ion 162 631
(116) (6241) (6280) (6100)	VEIF VEIF VRPI 6280 VERI	CIVONLSC RIVQMLAD 6 LISCNVCT	TIRGLSORY HL DLSDCV 290 KARYAYDSF	VENTANG VVLVTWAHO 6300 (TSYYGCW)	FELTCL Sect	RYFV ion 162 631
(116) (6241) (6280) (6100) (6668)	VEIF VRPE 6280 VORT	CIVOWLSO RIVOMLAD 6 LISCNVCT LOVESOG	Tirclserv HL Disdev 290 RFM AYDSP SCAPTENS!	VF VEWARD VULVTWARD 6300 (TGY VGCWI	FELTCL Sect THBYTCL KHCLCFL	KYFV RYFV ion 162 631 YLVIII
(116) (6241) (6280) (6100) (6668) (6184)	VEIF VRPE 6280 VCRE ISKS	CIVOWLSO RIVOMLAD 6 LISCNVCT 1-0V6S.G	Tirclserv HL Disdev 290 RM Aybsi Seartensi RM Cense	VFVEWAHO OLVTWAHO 6300 (TGYTGCW) LTOAYACWI LTOYTGCW)	TE OYSM SFELTCL Sect SHEYTCA KACLCFA RHSYSCA	K 1 - Vi RYFV ion 162
(116) (6241) (6280) (6100) (6668) (6184)	VEIF VRPE 6280 VCRE ISKS	CIVOWLSO RIVOMLAD 6 LISCNVCT 1-0V6S.G	TIRGLSORY HL DLSDCV 290 KARYAYDSF	VFVEWAHO OLVTWAHO 6300 (TGYTGCW) LTOAYACWI LTOYTGCW)	TE OYSM SFELTCL Sect SHEYTCA KACLCFA RHSYSCA	K 1 - Vi RYFV ion 162

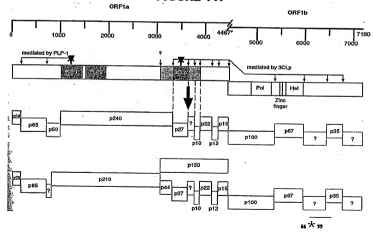
FIGURE 115 (contd.)

(6319)	6319		6330		6340		Section	6257
(6139)	TAVO	r comment	TESLSS	Marty CA	PCIELE	O B LIVE N	TO COMP	0337
(5706)	T. T. W.	rain the first	SSNAOP	trinting	re safete	TEN TIME	CAPP NO	OIL W.
(6223)	T. T. 17.7.7	されなかれば 1 mag 7 mag 2 を	TESLTS	14 12 13 13 13 13 13 13 13 13 13 13 13 13 13	0.01111	Tables Was	SATIST	OLK.
(104)			YE-PER	TA CF TVC. T	COVEN	Market San	DAL	DA. K
(6310)		TOOMON	SGSLSS	V.TELTE	TARXM	中面等同点	IASCDA	
(65 (6)	La Ji V Li.	TÖÜMET	2122722	DHELLE	CSVHK			
(00CO)	0050						 Section 	
(6358)	0358	ANTERNA	6370	स्टब्स्ट इस्ट इस्ट	6380_			6396
(61/8)	CLAV	ADCECK	NUMBER	EXPI	SNEES	INTSC	RATOR	VILL
(5/45)	CLAI	MNAFEC	DVUVLL	TYPHI	ANDDE	VNSSC	RYLOR	MYL
(6262)	CLAV	HOCECI	SWNWNL	EYPI	SNEVS	VNTSC	RLLOR	Val
(229)								
(6358)	CLAV	HDCFCN	VNWNL	EYPI	SNELS	VNTSC	CRLLQR	VML
							Section	165
(6397)	6397		,6410		6420			6435
(6217)	KAAM	LCNRYI	CCYDIC	NPKA	ACVED	EDE	KFYDA	OBT
(5784)	NACV	DALKVN	IVVYDIG	NPKG	KCVRR	GDVN	RFYDK	NPT
			VČÝĎÍC					
(229)			WARRIED TO THE SES		OFFICE CONTRACT		MANAGEMENT - 39-1960	
(6397)	KAAM	LCNRY	VCYDIG	NPKG	LACVK	FDF	KFYDA	NPT
							- Section	166
(6436)	6436		.64	50	.646	30		6474
(6254)	VKSV	KT LEES	FEARKE				Y PPN	Z VIXI
(5823)	VRNV	KOFEYI	YNQHKI	KEADO	TOMEN	NONVI	CABDA	STAV
(6338)	VKSV	KOEVYE	YEAHKI	OFTD	at/CM FM	NCNV	KYPAN	ΔWV
(229)		CHARLES TANKEN	***************************************		ALLEY STANISH AND AND AND AND AND AND AND AND AND AND	MODERNOON SONE WAS	SCHOOLSE TO	
		KOFLY	YEAHKI	F D(T.CMEN	MCNVI	א סציאר	AVV
(0.00)			TOMME	, , ,	JII CITE W		- Section	
(6475)	6475	6480	č	490		500	- Occion	6513
(0410)	2473	U-100	LÙLPGC	490	-21.14.1	OUU	T. Carle St. Ass. La	0013
(6283)	CHILD	THV LOT	Tribitation	NGUS	TA A IN REF	AT HT	KPESHA	AFE
(0002)	CKYL	TKHIJS	FINLPGC	Neep	DAAMKE	titi. XT.	KANA	SEE
(0000)	CELO	T. WATERS	Lint bec	NGGS	PAANKE	APHTI	PETRA	AFE
(229)								
(64/5)	CRFD	TRVLN	LNLPGO	NGGS	LYVNKI		PFSRA	
							Section	168
(6514)				6530		6540		6552
			SOTPC					
			POSSPCE					
			and the second second	Sec. of the second sec. of		4-14-4-		T man
(6416)	NIKP	WEEL	SDTPC	YMEG	MESKOI	DIAB	LRSAIC	LT
(229)			SDTPCY YSDTPCY					

FIGURE 115 (contd.)

	Section 169					
	6591	6580	6570	6560	6553	(6553)
	FWVYXTED	TAGET	EYLESYNTA	avcikharey	CNIGG	(6371)
	THE THE PERSON	用五 色形中	EFVITSYNAA	AVCKKHADMY	CULGG	(၁೪૩೪)
	FWVYKTED	TAGET	EYLESYNTA	AVCL KHAEEY.	CNLGG	(6455)
						(229)
	FWVYKTFD	TAGET	EYLESYNTA	AVCLKHAEEY	CNTCC	(6553)
	Section 170			67	2222	(0500)
	6630	6620	6610	6600	6592	(6592)
	SEMPCATE	IYTGQA	AAANTAKIE	NTETKLOSLE	T X MIN	(6410)
	SEMPTVIT	IYDAIA	TAYNMYKGG	KSESALOSED	PANTW	(58/6)
	SEEPCAVI	EDERA	VVYNEVNAC	UPETRICELE	E Y W TW	(229)
	~~~~~	VDC 2	WWYNTWEAC	NTFTKLOSLE		
	Section 171		VVINDVKAG	T		·····
	6669		6650	6640 Stop	6631	(6631)
	0009		THE SHIP OF THE PARTY	AKED KEDVVE		
	AND MEDICAL		UNOTET PER	vidogvek, vi	CDKVE	(6017)
	NAME OF THE PARTY	7767-6	KNNPLEDMA	aktonedvyvi	GEKVP	(6533)
	· · · · · · · · · · · · · · · · · · ·	STATE STATE	ericasia an annasia.		ATTEMPT TO THE STATE OF	(229)
	AKRSTR H	AVELE	INNTTEPTN	AKIQ EDVVV	GDKVI	(6631)
	Section 172					
	6708		6690	6680		(6670)
	LYGVCMYT	TECSO	HVIWOYARE	frnlnidvowi	PELKI.	(6488)
	LVKVCAYT	PIYRN	FVIWDYATE	LKGLCVDVTN)	PMNRI	(6056)
	LYKYCKYT	VECSS	HYLWDYAND	FRMLNIOVCK:	PELKL	(6572)
						(229)
	TYKVC YT	IFCSN	HVIWDYAKD	FRNLNIDVCW	PELKL	(6670)
	Section 173					
	6747		6730	6720		(6709)
	CSTTKVKS	MNCVY	ngaleafkr.	ÖKLNVL FÖĞRI	DEKEL	(6527)
	VSTQCYKR.	DMAVL	YCDYOSTLA	SLNVI, YDDI	DIEPN	(6095)
	LUTTKIKS	RNEVY	NGALEAFKE	eslavl pdgri	DIOCL	(6611)
						(229)
			NGALEAFKK.	D LNVLFDGR	Dr I	(6709)
	Section 174				0740	
	6786	12 CONTRACTO	6770	6760	6/48	(6748)
SEQ ID NO: 100	RKEGQDVI	FYFAV.	ARKAGDADC	GPPREEDNGV	LSMIK	(6400)
SEQ ID NO: 100	BUDAM	Allers Marie	PLKDG	GPPRAELNGV IPSHLLVQHGI GPOKADLNGV	YSYVE	(6732)
SEQ ID NO: 100	RKDGDDVI	FWTAV	AFKACD SDA	PROBADENGY	TRUTK	(0000)
SEQ ID NO: 999			WDKACDG2	GP RADLNGV		(229)
SEQ ID NO: 100	KKDGNDVI	r.w.r.AV	VDKVGDSD	GE KADINGV	POWIK	(0/40)





		7-7-				Section 1
(1) 1		10	20	30	40	51
(1) -	GIGHLE	CI NKEFS	CAAbs As Ad	TKALAATYU	THDEALVNV	EAGHERN
					TGDLAVELGI	
					/GGDL::VCLNV	
					/GEDI - VELNV	
(1)	SINTE	KDCSKSYS	SGYHPAHAPS	FLAVDDKYK	VGGDLAVCLNV	
						Section 2
(52) 5		.60	70	80	90	102
					RAWIGEDVEGO	
					(AWIGEDVESC	
					RAWIGPDVECO	
					RAWIGIOVECC	
(52)	KKLISM.	MGFKMNI	2VNGIPNMF1	TREEATRHVI	RAWIGFDVEGC	— Section 3
400) 4	100	440	.120	400	4.40	
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					RNNAOTSTSEO	
				DIEDNIKE	RVNAOTSTSEO	
					RVNAOTSTSEO	PRUITOIN
(103)	LUPEPO	LGESIGVI	NLVAVPIGIV	DIENNIKEL	KVNAQISISEQ	Section 4
(154)	154	160	.170	.180	.190	204
153)	YKGLEW	NVVRIKT	MOMES DILLIKE	IJS DRVVEVI	Wangfeltsmk	ургика сет
(153)	YKGLPW	NWVRTKT.	JOMESDTEKS	L'SDRVVFVL	WAHGEELTSMK	YEVKIGEE
(153)	YKGDEW	NVVRTRT	vomes de ekc	ESDRVVEVE	wahge elesik	ŶŦWKĪĠ₽Ŀ
(153)	KKGLEW	NYVETKI	VOMESDEEK	LSDRVVEVL	Wangfeltsmk	YEVKIGPI
(154)	YKGLPW	NVVRIKI	VQMLSDTLKG	LSDRVVFVL	WAHGFELTSMK	
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(204)	RICCIC	DKRATCE	STSSDTYAC	VNHSVGEDYV	YNPEMEDVOOW	GLYGNIQI
					YN P.BMT.DVQQW	
(204)	RICCLO	DEFATOR	STEEDTYAGE	inesverdyv	YNDFMIDVQQW	GLYGSLT
(204)	RICCLE	DKRATCE	STSSDTYACI	NHSVGFDYN	YNEFMIDVQQW	GLYCSLS
(205)	RTCCLC	DKRATCF	STSSDTYAC	NHSVGFDYV	YNPFMIDVQQW	
						Section 6
(256)	256		270	280	290	30
(255)	миртиз	H J M GH T- I	vasvoathii	CLAIMNAFE	CDVKWDLTZEH	TAREDEM
(255)	CHOTAC	STUKEAH	VASSDAIMTS	EXAVYDOR	nni aanvel ti	SELLSI
(255)	REDPIS	SVIIKGAN	VASSIMITE!	CLUANDE	RSVINNLETFI	TSHEVSV
(255)	PHOTHE	SWHKGAH	WASSUATET	STLEVHDOFO	nsvannleypi	TSHELSV
(256)	NHDLHO	SVHKGAH	VASSDAIMT	RCLAVHDCFC	NSVNWNLEYPI	ISNELSV

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## FIGURE 118 (contd.)

(307) 307	320	330	340	Section 7
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#### FIGURE 120

#### FIGURE 120A

PRHTQRT-PTVDSSL-WVSK-ITKSMVTLICLSPAKKLFVTFVRGLALM-RAVMQ LEMLWVLTYLSS-DFLQVLT--LYRLVMLTLKITQNSPELMQNLHQVTSLNILYH SCIKACPGM-CVLR-YKCSVIH-KDCQTESCSSFGRMALSLHQ-STLSRLDLKER VVCVTNVQLAFLLHQILMPAGIILWVLTMSITHL-LMFSSGALRVTFRVTMTNIA RYMEMHMWLVVMLS-LDV-QSMSALLSALIGLLNTLL-EMN-GLILLAEKYNTWL-SLHCLLISFQFFMT-EIQRLSSVCLRLK-NGSSTMLSHVVTKLTK-RNSSILML YITINSLMVFVCFGIVTLIVTQPMQLCVGLTQESCQT-TYQAVMVVVCM-ISMHS TLQLSIKVHLLI-SNCLSFTILIVLVSLMANK-CRILIMFHSNLLRVLHDAI-VV LFADTMOMSTDSTWMHII--FLLDLAYGFTNNLILITCGIHLPGYWV

#### FIGURE 120B

LGIPKGHDLP-THLYDGFQNELPSQWLP-YVYHPRRSYSSRSCVDWL-CRGLSCN-RCCGY-PTSPARIFYRC-LSSCTDWLC-H-K-HRIHQS-CKTSTR-PV-TSYTTHV-RLALECSAY-DSTNAQ-YTERIVRQSRVRPLGAWL-AYINEVLCQDWT-KNVLSV-QTCNLLFYFIRYLCLLESFCGF-LCL-PIYD-CSAVGLYG-PSE-P-PTLPGTWKCTCG-L-CYHD-MFSSP-VLC-AR-LVC-IPYYRR-TEG-FCLQKSTTHGCEVCIAC--VSSSS-HRKSKGYQVCASG-SRMEVLRCSAM-QSLQNRGTLLFLCYTSR-IH-WCLFVLEL-R-SLPSQCNCV-V-HKSLVKLELTRL-WW-FVCE-ACIPHSSFR-KCTY-FKAIAFLLLF-SL-VSWQTSSVGY-LCSTQICYVYYTMQFRWCCLQTPCK-VPTVLGCI-YDDFCWI-PMDLQTI-YL-PVEYIYOVTEF

#### FIGURE 120C

-AYPKDMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVG
TNLPLQLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPMNVV
RIKIVQMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSD
TYACWNHSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAV
HECFVKRVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVP
QAEVEWKFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFD
TRVLSNLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSD
LDYVPLKSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNT
FTRLQSL

#### FIGURE 120D

-TL-PGKCIPQVISIKLFVNP-AKSSRNHHIICIQVLSVLICMVSANSTT-IASCNTRSR FEWNIINIRHYLFAMRLTRTIRIVKERQLL-ISKCTFIESWSVECMLIHIQTTTITAW-V QV-QDSCVKPTHNCIGWVTINVTIPKQTNTISEFIVMYSIRIEEFLYFVSFVTTWLSIVE LPFYFSLRHTLDSLWISYVMKNWKLISKQCRLHNHVLYFSASRINPQFISYNRVFNRPIN ALNKALMDC-TSSHDSITTSIMCISMYLAMLVMVTLKVTRKAPLLNINHKWVIDIVKTHR MIPAGISI-SRKASCTFVTQTTRSFRSNLDKVLH-CKLKAMRPKDEHDSV-QSFQCITE-HLYYLNTHYIPGQAFIHEWYKMFKLVTWWRFCINSGEFCVIFSVNITSRYSY-VNTCRKS-LER-VSTHSISSCMTALYIKANPRTNVTNSFFAGDKHIRVTIDLVIHFETHHRDESTVGHVLWVCL

#### FIGURE 120E

KLCNLVNVFHRL-VSNCL-IHRLNPAEIIILYASKYCRYSFAWCLQTAPPKLHRVIHVAD LSGT-SISDTTCLP-DSQGLSE--KKGNCFKLVNALLSKAGVWNACLFTYKLPPSQPGKF KFDKTLVSNLHTIALAG-RSTLQFQNKQTPSVNLS-CIA-E-KSSSIL-ALSLHG-AS-N FHSTSA-GTHLIAFGFPMS-RTGNLSASNADFTTMCCTFLQAELTLSSSPIIGYSTDQST RLTKHSWTAKHLVMIASQLATCAFPCTWQCWSWLL-RLPVKPHC-TSIINGL-T-SKPTE-FQQA-VSDEVEKQVARLSHRQHVLSGPILTTKYFIDVSSKPCAQRTNTTLSDNPFSVSLS ICTILIRTTFQGKPLYMSGIRCLNWSPGGGFALTLVNSVLFSVST-PVGTATKLTPVENPSWRGRLVPTASLVA-QPSTSKPIHART-RIASSRVINILG-PLTW-FILKPIIEMSLR-V MSFGYA-

#### FIGURE 120F

NSVTW-MYSTGYKYQIVCKSIG-IQQKSSYYMHPSTVGTHLHGVCKQHHLNCIV-YT-QI
-VEHNQYPTLLVCHETHKDYQNSKRKAIALN--MHFYRKLECGMHAYSHTNYHHHSLVSS
SLTRLLCQTYTQLHWLGNDQRYNSKTNKHHQ-IYRDV-HKNRRVPLFCKLCHYMAEHRRT
SILLQEEAHT--PLDFLCHEELETYQQAMQTSQPCVVLFCKQN-PSVHLL--GIQQTNQR
A-QSTHGLLNI-S--HHN-PHVHFHVPGNVGHGYSEGYP-SPTAEHQS-MGYRHSQNPQN
DSSRHKYLMK-KSKLHVCHTDNTFFQVQS-QSTSLM-AQSHAPKGRTRLCLTILSVYH-A
FVLS-YALHSRASLYT-VV-DV-TGHLVEVLH-LW-ILCYFQCQHNQSVQLLS-HL-KIL
AGEVG-YPQHL-LHDSPLHQSQSTHERDE-LLRG--TY-GNH-LGNSF-NPS-R-VYGRS
CPLGMPR

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## FIGURE 121

	10	20	30	40	50	60
SEQ ID NO:10033		<u>'</u> -			I CTCTATGATGG	
SEQ ID NO:10084	CCTAGGCATAC	CCAAAGGACA				
Consensus Prim. cons.	CCTAGGCATAC	~~~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~			CTCTATGATGG	
FIIM. COMS.	CCIAGGCAIAC	CAMMOGACA	TIGACCIACC	SIAGACICAI	LICIAIGAIGG	GITICAAAA
	70 1	80 I	90 I	100	110	120
SEQ ID NO:10033	TGAATTACCAA	GTCAATGGTT	ACCCTAATAT	GTTTATCAC	CCGCGAAGAAG	CTATTCGTC
SEQ ID NO:10084						
Consensus	TGAATTACCAA					
Prim cons.	TGAATTACCAA	GTCAATGGTI	ACCCTAATA	PGTTTATCAC(	CCGCGAAGAAG	CTATTCGTC
	130	140	150	160	170	180
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SEQ ID NO:10033	ACGTTCGTGCG	TGGATTGGĊI	TTGATGTAĠ	AGGGCTGTCA	IGCAACTAGAG	ATGCTGTGG
SEQ ID NO:10084						
Consensus	ACGTTCGTGCG'					
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SEQ ID NO:10033						
SEQ ID NO:10084	GTACTAACCTAG					
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Prim. cons.	CTGGTTATGTT					
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Consensus	CAGGTGACCAG			CAIGIAIAA	MGGC11GCCC1	GALDIAND
Prim. cons.	CAGGTGACCAG			ICATGTATAA.	AGGCTTGCCCT	GGAATGTAG

etc.

#### FIGURE 122

#### 5'3' Frame 1

cctaggcatacccaaaggacatgacctaccgtagactcatctctatgatgggtttcaaaa PRHTORT-PTVDSSL-WVSK tgaattaccaagtcaatggttaccctaatatgtttatcacccgcgaagaagctattcgtc - I T K S M V T L I C L S P A K K L F V  ${\tt acgttcgtgcgtggattggctttgatgtagagggctgtcatgcaactagagatgctgtgg}$ T F V R G L A L M - R A V M Q L E M L W gtactaacctacctctccagctaggattttctacaggtgttaacttagtagctgtaccga V L T Y L S S - D F L Q V L T - - L Y R ctggttatgttgacactgaaaataacacagaattcaccagagttaatgcaaaacctccac LVMLTLKITQNSPELMQNLH caggtgaccagtttaaacatcttataccactcatgtataaaggcttgccctggaatgtag QVTSLNILYHSCIKACPGM-C V L R - Y K C S V I H - K D C Q T E S tgttcgtcctttgggcgcatggctttgagcttacatcaatgaagtactttgtcaagattg C S S F G R M A L S L H Q - S T L S R L gacctgaaagaacgtgttgtctgtgtgacaaacgtgcaacttgcttttctacttcatcag DLKERVVCVTNVQLAFLLHQ atacttatgcctgctggaatcattctgtgggttttgactatgtctataacccatttatga ILMPAGIILWVLTMSITHL ttgatgttcagcagtggggctttacgggtaaccttcagagtaaccatgaccaacattgcc LMFSSGALRVTFRVTMTNIA aggtacatggaaatgcacatgtggctagttgtgatgctatcatgactagatgtttagcag RYMEMHMWLVVMLS-LDV-Q tccatgagtgctttgttaagcgcgttgattggtctgttgaataccctattataggagatg S M S A L L S A L I G L L N T L L - E M aactgagggttaattctgcttgcagaaaagtacaacacatggttgtgaagtctgcattgc N - G L I L A E K Y N T W L - S L H C ttgctgataagtttccagttcttcatgacataggaaatccaaaggctatcaagtgtgtgc L I S F Q F F M T - E I Q R L S S V C  $\verb|ctcaggctgaagtagaatggaagttctacgatgctcagccatgtagtgacaaagcttaca|\\$ LRLK-NGSSTMLSHVVTKLT aaatagaggaactcttctattcttatgctatacatcacgataaattcactgatggtgtttK - R N S S I L M L Y I T I N S L M V F gtttgttttggaattgtaacgttgatcgttacccagccaatgcaattgtgtgtaggtttg V C F G I V T L I V T Q P M Q L C V G L acacaagagtcttgtcaaacttgaacttaccaggctgtgatggtggtagtttgtatgtga TQESCQT-TYQAVMVVCMataagcatgcattccacactccagctttcgataaaagtgcatttactaatttaaagcaat ISMHSTLQLSIKVHLLI-SN tgcctttcttttactattctgatagtccttgtgagtctcatggcaaacaagtagtgtcggC L S F T I L I V L V S L M A N K - C R atattgattatgttccactcaaatctgctacgtgtattacacgatgcaatttaggtggtg I L I M F H S N L L R V L H D A I - V V ctgtttgcagacaccatgcaaatgagtaccgacagtacttggatgcatataatatgatga L F A D T M Q M S T D S T W M H I I - tttctgctggatttagcctatggatttacaaacaatttgatacttataacctgtggaata F L L D L A Y G F T N N L I L I T C G I

catttaccaggttacagagttta H L P G Y R V

#### 5'3' Frame 2

cctaggcatacccaaaggacatgacctaccgtagactcatctctatgatgggtttcaaaat LGIPKGHDLP-THLYDGFON gaattaccaagtcaatggttaccctaatatgtttatcacccgcgaagaagctattcgtca ELPSOWLP-YVYHPRRSYS cqttcqtqcqtqqattqqctttqatqtaqaqqqctqtcatqcaactaqaqatqctqtqqq R S C V D W L - C R G L S C N - R C C G tactaacctacctctccagctaggattttctacaggtgttaacttagtagctgtaccgac Y - P T S P A R I F Y R C - L S S C T D tggttatgttgacactgaaaataacacagaattcaccagagttaatgcaaaacctccacc W L C - H - K - H R I H O S - C K T S T aggtgaccagtttaaacatcttataccactcatgtataaaggcttgccctggaatgtagt R - P V - T S Y T T H V - R L A L E C S AY-DSTNAQ-YTERIVRQSR gttcgtcctttgggcgcatggctttgagcttacatcaatgaagtactttgtcaagattgg V R P L G A W L - A Y I N E V L C Q D W acctgaaagaacgtgttgtctgtgtgacaaacgtgcaacttgcttttctacttcatcaga T - K N V L S V - Q T C N L L F Y F I R tacttatgcctgctggaatcattctgtgggtttttgactatgtctataacccatttatgat Y L C L L E S F C G F - L C L - P I Y D tgatgttcagcagtggggctttacgggtaaccttcagagtaaccatgaccaacattgcca - C S A V G L Y G - P S E - P - P T L P ggtacatggaaatgcacatgtggctagttgtgatgctatcatgactagatgtttagcagt G T W K C T C G - L - C Y H D - M F S S ccatgagtgctttgttaagcgcgttgattggtctgttgaataccctattataggagatga P-VLC-AR-LVC-IPYYRRactgagggttaattctgcttgcagaaaagtacaacacatggttgtgaagtctgcattgct TEG-FCLQKSTTHGCEVCIA tgctgataagtttccagttcttcatgacataggaaatccaaaggctatcaagtgtgtgcc C - - V S S S S - H R K S K G Y O V C A tcaggctgaagtagaatggaagttctacgatgctcagccatgtagtgacaaagcttacaa SG-SRMEVLRCSAM--OSLO aatagaggaactcttctattcttatgctatacatcacgataaattcactgatggtgtttg NRGTLLFLCYTSR-IH-WCL tttgttttggaattgtaacgttgatcgttacccagccaatgcaattgtgtgtaggtttga FVLEL-R-SLPSQCNCV-Vcacaagagtcttgtcaaacttgaacttaccaggctgtgatggtggtagtttgtatgtgaa H K S L V K L E L T R L - W W - F V C E taagcatgcattccacactccagctttcgataaaagtgcatttactaatttaaagcaatt - A C I P H S S F R - K C I Y - F K A I gcctttcttttactattctgatagtccttgtgagtctcatggcaaacaagtagtgtcgga tattgattatgttccactcaaatctgctacgtgtattacacgatgcaatttaggtggtgc Y - L C S T Q I C Y V Y Y T M Q F R W C totttgcagacaccatgcaaatgagtaccgacagtacttggatgcatataatatgatgat

C L Q T P C K - V P T V L G C I - Y D D ttctgctggatttagcctatggatttacaaacaatttgatacttataacctgtggaatac F C W I - P M D L Q T I - Y L - P V E Y atttaccaggttacagagttta I Y O V T E F

#### 5'3' Frame 3

cctaggcatacccaaaggacATGacctaccgtagactcatctctatgatgggtttcaaaatg - A Y P K D M T Y R R L I S M M G F K M aattaccaagtcaatggttaccctaatatgtttatcacccgcgaagaagctattcgtcac NYQVNGYPNMFITREEAIRH gttcgtgcgtggattggctttgatgtagagggctgtcatgcaactagagatgctgtgggt V R A W I G F D V E G C H A T R D A V G actaacctacctctccagctaggattttctacaggtgttaacttagtagctgtaccgact TNLPLQLGFSTGVNLVAVPT ggttatgttgacactgaaaataacacagaattcaccagagttaatgcaaaacctccacca G Y V D T E N N T E F T R V N A K P P P ggtgaccagtttaaacatcttataccactcatgtataaaggcttgccctggaatgtagtg G D Q F K H L I P L M Y K G L P W N V V RIKIVQMLS'DTLKGLSDRVV ttcgtcctttgggcgcatggctttgagcttacatcaatgaagtactttgtcaagattgga F V L W A H G F E L T S M K Y F V K I G cctgaaagaacgtgttgtctgtgtgacaaacgtgcaacttgcttttctacttcatcagat PERTCCLCDKRATCFSTSSD acttatgcctgctggaatcattctgtgggttttgactatgtctataacccatttatgatt TYACWNHSVGFDYVYNPFMI gatgttcagcagtggggctttacgggtaaccttcagagtaaccatgaccaacattgccag D V Q Q W G F T G N L Q S N H D O H C O gtacatggaaatgcacatgtggctagttgtgatgctatcatgactagatgtttagcagtc V H G N A H V A S C D A I M T R C L A V catgagtgctttgttaagcgcgttgattggtctgttgaataccctattataggagatgaa HECFVKRVDWSVEYPIIGDE ctgagggttaattctgcttgcagaaaagtacaacacatggttgtgaagtctgcattgctt L R V N S A C R K V Q H M V V K S A L L gctgataagtttccagttcttcatgacataggaaatccaaaggctatcaagtgtgtgcct ADKFPVLHDIGNPKAIKCVP caggctgaagtagaatggaagttctacgatgctcagccatgtagtgacaaagcttacaaa Q A E V E W K F Y D A Q P C S D K A Y K atagaggaactcttctattcttatgctatacatcacgataaattcactgatggtgtttgt I E E L F Y S Y A I H H D K F T D G V C ttgttttggaattgtaacgttgatcgttacccagccaatgcaattgtgtgtaggtttgac LFWNCNVDRYPANAIVCRFD acaagagtcttgtcaaacttgaacttaccaggctgtgatggtggtagtttgtatgtgaat TRVLSNLNLPGCDGGSLYVN aagcatgcattccacactccagctttcgataaaagtgcatttactaatttaaagcaattg K H A F H T P A F D K S A F T N L K O L cctttcttttactattctgatagtccttgtgagtctcatggcaaacaagtagtgtcggat P F F Y Y S D S P C E S H G K Q V V S D

#### 3'5' Frame 1

- T L - P G K C I P Q V I S I K L F V N ccataggctaaatccagcagaaatcatcatattatatgcatccaagtactgtcggtactc P-AKSSRNHHIICIOVLSVL atttqcatqqtqtctgcaaacagcaccacctaaattqcatcqtqtaatacacqtaqcaqa ICM-VSANSTT-IASCNTRSR tttgagtggaacataatcaatatccgacactacttgtttgccatgagactcacaaggact F E W N I I N I R H Y L F A M R L T R T atcagaatagtaaaagaaaggcaattgctttaaattagtaaatgcacttttatcgaaagc IRIVKERQLL - ISKCTFIES tggagtgtggaatgcatgcttattcacatacaaactaccaccatcacagcctggtaagtt W S V E C M L I H I Q T T T I T A W - V caaqtttgacaagactcttgtgtcaaacctacacacaattgcattggctgggtaacgatc O V - O D S C V K P T H N C I G W V T T aacqttacaattccaaacaacaacaccatcaqtqaatttatcqtqatqtataqcata NVTIPKQTNTISEFIVMYSI agaatagaagagttcctctattttgtaagctttgtcactacatggctgagcatcgtagaa RIEEFLYFVSFVTTWLSIVE cttccattctacttcagcctgaggcacacacttgatagcctttggatttcctatgtcatg L P F Y F S L R H T L D S L W I S Y V M aagaactggaaacttatcagcaagcaatgcagacttcacaaccatgtgttgtacttttct K N W K L I S K O C R L H N H V L Y F S gcaagcagaattaaccctcagttcatctcctataatagggtattcaacagaccaatcaac A S R I N P O F I S Y N R V F N R P I N gcgcttaacaaagcactcatggactgctaaacatctagtcatgatagcatcacaactagc ALNKALMDC - TSSHDSITTS cacatgtgcatttccatgtacctggcaatgttggtcatggttactctgaaggttacccgt H M C I S M Y L A M L V M V T L K V T R aaaqccccactqctqaacatcaatcataaatqqqttataqacataqtcaaaacccacaqa K A P L L N I N H K W V I D I V K T H R atgattccagcaggcataagtatctgatgaagtagaaaagcaagttgcacgtttqtcaca M I P A G I S I - - S R K A S C T F V T cagacaacacgttctttcaggtccaatcttgacaaagtacttcattgatgtaagctcaaa O T T R S F R S N L D K V L H - C K L K gccatgcgcccaaaggacgaacacgactctgtctgacaatcctttcagtgtatcactgag AMRPKDEHDSV-QSFQCITE catttgtactatcttaatacgcactacattccagggcaagcctttatacatgagtggtat H L Y Y L N T H Y I P G Q A F I H E W Y aagatgtttaaactggtcacctggtggaggttttgcattaactctggtgaattctgtgtt K M F K L V T W W R F C I N S G E F C V attiticate transcrapt contact and transcrapt conta

#### 3'5' Frame 2

K L C N L V N V F H R L - V S N C L - I cataggetaaatecageagaaateateatattatatgeateeaagtaetgteggtaetea H R L N P A E I I I L Y A S K Y C R Y S tttgcatggtgtctgcaaacagcaccacctaaattgcatcgtgtaatacacgtagcagat FAWCLOTAPPKLHRVIHVAD ttgagtggaacataatcaatatccgacactacttgtttgccatgagactcacaaggacta LSGT-SISDTTCLP-DSQGL tcagaatagtaaaaggaaaggcaattgctttaaattagtaaatgcacttttatcgaaagct SE--KKGNCFKLVNALLSKA ggagtgtggaatgcatgcttattcacatacaaactaccaccatcacagcctggtaagttcG V W N A C L F T Y K L P P S Q P G K F  ${\tt aagtttgacaagactcttgtgtcaaacctacacacaattgcattggctgggtaacgatca}$ K F D K T L V S N L H T I A L A G - R S acgttacaattccaaaacaacaacaacatcagtgaatttatcgtgatgtatagcataa TLQFQNKQTPSVNLS-CIAgaatagaagagttcctctattttgtaagctttgtcactacatggctgagcatcgtagaac E - K S S S I L - A L S L H G - A S - N ttccattctacttcagcctgaggcacacacttgatagcctttggatttcctatgtcatga FHSTSA-GTHLIAFGFPMSagaactggaaacttatcagcaagcaatgcagacttcacaaccatgtgttgtacttttctg RTGNLSASNADFTTMCCTFL caagcagaattaaccctcagttcatctcctataatagggtattcaacagaccaatcaacg Q A E L T L S S S P I I G Y S T D Q S T cgcttaacaaagcactcatggactgctaaacatctagtcatgatagcatcacaactagcc R L T K H S W T A K H L V M I A S Q L A acatgtgcatttccatgtacctggcaatgttggtcatggttactctgaaggttacccgta T C A F P C T. W Q C W S W L L - R L P V aagccccactgctgaacatcaatcataaatgggttatagacatagtcaaaacccacagaa KPHC-TSIINGL-T-SKPTE tgattccagcaggcataagtatctgatgaagtagaaaagcaagttgcacgtttgtcacac - F Q Q A - V S D E V E K Q V A R L S H agacaacacgttctttcaggtccaatcttgacaaagtacttcattgatgtaagctcaaag RQHVLSGPILTKYFIDVSSK ccatgcgcccaaaggacgaacacgactctgtctgacaatcctttcagtgtatcactgagc P C A Q R T N T T L S D N P F S V S L S

#### 3'5' Frame 3

NSVTW-MYSTGYKYQIVCKS ataggctaaatccagcagaaatcatcatattatatgcatccaagtactgtcggtactcat IG-IOOKSSYYMHPSTVGTH ttgcatggtgtctgcaaacagcaccacctaaattgcatcgtgtaatacacgtagcagatt L H G V C K Q H H L N C I V - Y T - Q I tgagtggaacataatcaatatccgacactacttgtttgccatgagactcacaaggactat - V E H N Q Y P T L L V C H E T H K D Y cagaatagtaaaaggaaatgctttaaattagtaaatgcacttttatcgaaagctg Q N S K R K A I A L N - - M H F Y R K L gagtgtggaatgcatgcttattcacatacaaactaccaccatcacagcctggtaagttca E C G M H A Y S H T N Y H H H S L V S S agtttgacaagactcttgtgtcaaacctacacacaattgcattggctgggtaacgatcaa S L T R L L C Q T Y T O L H W L G N D O cgttacaattccaaaacaaacaaccatcagtgaatttatcgtgatgtatagcataag RYNSKTNKHHO-IYRDV-HK aatagaagagttcctctattttgtaagctttgtcactacatggctgagcatcgtagaact NRRVPLFCKLCHYMAEHRRT tccattctacttcagcctgaggcacacacttgatagcctttggatttcctatgtcatgaa SILLQPEAHT--PLDFLCHE gaactggaaacttatcagcaagcaatgcagacttcacaaccatgtgttgtacttttctgc E L E T Y Q Q A M Q T S Q P C V V L F C aagcagaattaaccctcagttcatctcctataatagggtattcaacagaccaatcaacgc KON-PSVHLL--GIOOTNOR gcttaacaaagcactcatggactgctaaacatctagtcatgatagcatcacaactagcca A - O S T H G L L N I - S - - H H N - P catgtgcatttccatgtacctggcaatgttggtcatggttactctgaaggttacccgtaa HVHFHVPGNVGHGYSEGYPagccccactgctgaacatcaatcataaatgggttatagacatagtcaaaacccacagaat SPTAEHQS-MGYRHSQNPQN gattccagcaggcataagtatctgatgaagtagaaaagcaagttgcacgtttgtcacaca D S S R H K Y L M K - K S K L H V C H T gacaacacgttctttcaggtccaatcttgacaaagtacttcattgatgtaagctcaaagc

D N T F F Q V Q S - Q S T S L M - A O S catqcqcccaaaggacgaacacgactctgtctqacaatcctttcagtgtatcactgagca HAPKGRTRLCLTILSVYH-A tttqtactatcttaatacgcactacattccagggcaagcctttatacatgagtggtataa FVLS-YALHSRASLYT-VVgatgtttaaactggtcacctggtggaggttttgcattaactctggtgaattctgtgttat DV-TGHLVEVLH-LW-ILCY tttcagtgtcaacataaccagtcggtacagctactaagttaacacctgtagaaaatccta FOCQHNQSVQLLS-HL-KI-L gctggagaggtaggttagtacccacagcatctctagttgcatgacagccctctacatcaa AGEVG-YPQHL-LHDSPLHO agccaatccacgcacgaacgtgacgaatagcttcttcgcgggtgataaacatattagggt SQSTHERDE-LLRG--TY-G aaccattgacttggtaattcattttgaaacccatcatagagatgagtctacggtaggtca NH-LGNSF-NPS-R-VYGRS tatcctttgggtatgcctagg CPLGMPR

# FIGURE 123

CCTAGGCATACCCAAAGGACATGACCTACCGTAGACTCATCTCTATGATGGGTTCAAAAATGAATTACCAAGTCAATGGT
i
${\tt TACCCTAATATGTTTATCACCCGCGAAGAAGCTATTCGTCACGTTCGTGCGTG$
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$
NCTGGTTATGTTGACACTGAAAATAACACAGAATTCACCAGAGTTAATGCAAAACCTCCACCAGGTGACCAGTTTAAACAC
CTGGTTATGTTGACACTGAAAATAACACAGAATTCACCAGAGTTAATGCAAAACCTCCACCAGGTGACCAGTTTAAACATNN.
. N
CTTATACCACTCATGTATAAAGGCTIGCCCTGGAATGTGGGTATTAAGATAGTACAAATGCTCAGTGATACACTGAA
AGGATTGTCAGACAGAGTCGTGTTCGTCCTTTGGGCGCATGGCTTTGAGCTTACATCAATGAAGTACTTTGTCAAGATTG
N
GACCTGAAAGAACGTGTTGTCTGTGTGACAAACGTGCAACTTGCTTTTCTACTTCATCAGATACTTATGCCTGCTGGAAT
CATTCTGTGGGTTTTGACTATGTCTATAACCCATTTATGATTGAT
${\tt TAACCATGACCAACATTGCCAGGTACATGGAAATGCACATGTGGCTAGTTGTGATGCTATCATGACTAGATGTTTAGCAG}$
NNNN
${\tt TCCATGAGTGCTTTGATTAGGGGGTTAATTCTGCT}$
N
${\tt TGCAGAAAAGTACAACACATGGTTGTGAAGTCTGCATTGCTTGATAAGTTTCCAGTTCTTCATGACATAGGAAATCC}$
ii
${\tt AAAGGCTATCAAGTGTGTGCCTCAGGCTGAAGTAGAATGGAAGTTCTACGATGCTCAGCCATGTTAGTGACAAAGCTTACA}$
AAATAGAGGAACTCTTCTATTCTTATGCTATACATCACGATAAATTCACTGATGGTGTTTGTT
GTTGATCGTTACCCAGCCAATGCAATTGTGTGTGTGGGTTTGACACAAGAGTCTTGTCAAACTTGAACTTACCAGGCTGTGA
NN.
TGGTGGTAGTTTGTATGTGAATAAGCATGCATTCCACACTCCAGCTTTCGATAAAAGTGCATTTACTAATTTAAAGCAAT
NN.
${\tt TGCCTTTCTTTTACTATTCTGATAGTCCTTGTGAGTCTCATGGCAAACAAGTAGTGTCGGATATTGATTATGTTCCACTC}$
iii
A A A T C T G C T G T A C A C G A T T A C G A C T G C A C A T G G G C A C T G C A A T G G G C A C T C C A C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C C A C C C A C C C C C C C C
NNNNNN
${\tt GGATGCATATAATATGATGATTTCTGCTGGATTTAGCCTATGGATTTACAAACAA$
N
CAMMINACIA COMPACIA CAMMINA MACA TO MACA TO MACA

# FIGURE 123 (contd.)

ontu.,		
Pos	Score	Pred
21	0.651	 
45	0.354	Yes
48	0.387	_
60	0.590	Yes
76	0.470	_
90	0.676	Yes
145	0.192	_
160	0.410	-
172 247	0.290	_
286	0.221 0.219	_
333	0.373	_
355	0.178	_
381	0.286	_
439	0.405	-
459	0.204	_
547	0.289	_
580 597	0.447 0.449	_
604	0.290	_
646	0.427	_
667	0.427	_
673	0.208	
679	0.317	
694	0.180	
702 710	0.554 0.151	Yes
724	0.151 0.384	_
778	0.151	_
819	0.711	Yes
865	0.306	_
917	0.230	_
931	0.214	_
9 <b>41</b> 985	0.190	_
1012	0.274 0.368	_ _ _ _
1060	0.206	_
1120	0.193	_
1135	0.185	_
1147	0.431	_
1240	0.562	Yes
1270	0.377	_
1304 1336	0.190 0.353	-
1342	0.353	
1363	0.213	_
1374	0.178	_
1377	0.096	_
1400	0.056	_

# FIGURE 124

Sequences:	*	(bits)	Value
gi   74827   p gi   1491704 gi   2600754 gi   7769342 gi   6625761 gi   2641128 gi   4377413 gi   133592   gi   2600808 gi   1507782 gi   1803397 gi   7769353 gi   2512157 gi   2600809 gi   124246 gi   1414903 gi   458735   gi   2929345 gi   2929345 gi   2929345 gi   1938758 gi   1938758 gi   133591   gi   133591	ir   VFIHJH genome polyprotein 1b - murine hepatit  4   sp   P29982   RRPB_CVMJH RNA-directed RNA polymeras  6   ref   NP_068668.2   ORF1ab polyprotein [Murine hep  gb   AAF69332.1   AF208066_2 RNA-directed RNA polyme  gb   AAB86818.1   RNA-directed RNA polymerase [muri  emb   CAA36202.1   open reading frame 1b (AA 1-2733  sp   P16342   RRPB_CVMA5 RNA-DIRECTED RNA POLYMERASE  0   ref   NP_150073.2   orf1ab polyprotein [Bovine cor  0   gb   AAK83365.1   replicase [bovine coronavirus]  2   gb   AAL57305.1   replicase [bovine coronavirus]  2   gb   AAL57305.1   replicase [bovine coronavirus]  2   gb   AAL40397.1   AF220295_2 RNA polymerase 1b [bov  1   ref   NP_740140.1   coronavirus nspl1 [Murine hepa  2   ref   NP_742140.1   oronavirus nspl1 [Bovine coro]  3   emb   CAC39112.1   replicase polyprotein   favian  sp   P26314   RRPB_IBUS RNA-DIRECTED RNA POLYMERASE (  4   gb   AAO67706.1   ORF1ab polyprotein [Avian infecti  pref   NP_058422.1   replicase   Transmissible   gastro  2   ref   NP_058422.1   replicase   Transmissible   gastro  2   ref   NP_073549.1   replicase   polyprotein 1 ab   flum  sp   P18458   RRPB_BEV RNA-directed RNA polymerase (O    db   BAA13323.1   cyanoprotein alpha subunit precu	638 637 637 637 637 635 634 633 633 633 622 617 575 570 565 559 545 541 535 35	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0
(strain JH	pir  VFIHJH genome polyprotein 1b - murine hepatitis M) Length = 2731	virus	
	638 bits (1645), Expect = 0.0 s = 287/481 (59%), Positives = 366/481 (76%), Gaps = 5,	/481 (18	;)
Query: 6 Sbjct: 158	MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAV +TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HATRD++ 5 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHATRDSJ	GTN PLC	2
Query: 66	LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNV		
Sbjct: 164	LGFSTG++ V TG + F + A+ PPG+QFKHL+PLM +G W+\ 5 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEQFKHLVPLMSRGQKWD\	/VRI+IVÇ /VRIRIVÇ	
Query: 126	MLSD L L+D VV V WA FELT ++YF K+G E C +C+KRATCF++ +	Y CW	
Sbjct: 170	5 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGKEVVCSVCNKRATCFNSR		1764
Query: 186	HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLA HS DY+YNP++D+QOWG+TG+LSNHDCVHAHVASDAIMTRCLA		
Sbjct: 176	5 HSYSCDYLYNPLIVDIQQWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLA		

Ouerv: 246 RVDWSVEYPIIGDELRVNSACRKVOHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW 305 V+W++EYPII +E+ VN++CR +O ++ ++A+L +++ V +DIGNPK + CV Sbict: 1825 SVNWNLEYPIISNEVSVNTSCRLLORVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1882 Ouerv: 306 KFYDAOPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVLS Sbjct: 1883 KFYDASPV---VKSVKOFVYKYEAHKDOFLDGLCMFWNCNVDKYPANAVVCRFDTRVLSK 1939 Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC Sbjct: 1940 LNLPGCNGGSLYVNKHAFHTNPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 1999 Ouerv: 426 KSATCITRCNLGGAVCRHHANEYROYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLOS 485 +SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS Sbict: 2000 RSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLOS 2059 Query: 486 L 486 Sbict: 2060 L 2060 >gi|14917044|sp|P29982|RRPB_CVMJH RNA-directed RNA polymerase (ORF1B) gi | 7583321 | gb | AAA46458.2 | open reading frame 1b [murine hepatitis virus] Length = 2731Score = 637 bits (1644), Expect = 0.0 Identities = 287/481 (59%), Positives = 366/481 (76%), Gaps = 5/481 (1%) Ouerv: 6 MTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65 +TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HATRD++GTN PLO Sbjct: 1585 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHATRDSIGTNFPLO 1644 Ouery: 66 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHLIPLMYKGLPWNVVRIKIVO 125 LGFSTG++ V TG + F + A+ PPG+OFKHL+PLM +G W+VVRI+IVO Sbjct: 1645 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEQFKHLVPLMSRGQKWDVVRIRIVQ 1704 Ouerv: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185 MLSD L L+D VV V WA FELT ++YF K+G E C +C+KRATCF++ + Y CW Sbjct: 1705 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGKEVVCSVCNKRATCFNSRTGYYGCWR 1764 Query: 186 HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245 DY+YNP ++D+QQWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K Sbjct: 1765 HSYSCDYLYNPLIVDIQQWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK 1824 Query: 246 RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPOAEVEW 305 V+W++EYPII +E+ VN++CR +Q ++ ++A+L +++ V +DIGNPK + CV Sbjct: 1825 SVNWNLEYPIISNEVSVNTSCRLLQRVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1882 Query: 306 KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVLS Sbjct: 1883 KFYDASPV---VKSVKQFVYKYEAHKDQFLDGLCMFWNCNVDKYPANAVVCRFDTRVLSK 1939 Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC Sbjct: 1940 LNLPGCNGGSLYVNKHAFHTNPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 1999

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Ouerv: 426 KSATCITRCNLGGAVCRHHANEYROYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLOS 485
            +SATCITRCNLGGAVC HA EYR+YL++YN
                                           +AGF+ W+YK FD YNLWNTFTRLOS
Sbict: 2000 RSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLOS 2059
Ouerv: 486 L 486
Sbict: 2060 L 2060
>gi|26007546|ref|NP_068668.2| ORF1ab polyprotein [Murine hepatitis virus]
         Length = 7178
 Score = 637 bits (1644), Expect = 0.0
 Identities = 286/481 (59%), Positives = 364/481 (75%), Gaps = 5/481 (1%)
            MTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65
Ouerv: 6
            +TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HA RD++GTN PLQ
Sbjct: 6032 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHAIRDSIGTNFPLO 6091
            LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125
Ouery: 66
            LGFSTG++ V TG + F + A+ PPG+QFKHLIPLM +G W+VVRI+IVQ
Sbjct: 6092 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEQFKHLIPLMSRGQKWDVVRIRIVQ 6151
Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185
            MLSD L L+D VV V WA FELT ++YF K+G E C +C KRATCF++ + Y CW
Sbjct: 6152 MLSDHLADLADSVVLVTWAASFELTCLRYFAKVGREVVCSVCTKRATCFNSRTGYYGCWR 6211
Ouery: 186 HSVGFDYVYNPFMIDVOOWGFTGNLOSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245
            HS DY+YNP ++D+OOWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K
Sbjct: 6212 HSYSCDYLYNPLIVDIQOWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK 6271
Ouery: 246 RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW 305
             V+W++EYPII +E+ VN++CR +O ++ ++A+L +++ V +DIGNPK + CV
 Sbict: 6272 SVNWNLEYPIISNEVSVNTSCRLLQRVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 6329
Ouery: 306 KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365
                        +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+
            KEYDA P
 Sbjct: 6330 KFYDASPV---VKSVKQFVYKYEAHKDQFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 6386
Ouery: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKOVVSDIDYVPL 425
                                                                  +DYVPL
            LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC
 Sbjct: 6387 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 6446
 Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQS 485
            +SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLQS
 Sbjct: 6447 RSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLQS 6506
 Query: 486 L 486
 Sbict: 6507 L 6507
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>gi|7769342|gb|AAF69332.1|AF208066_2 RNA-directed RNA polymerase [murine hepatitis virus] Leneth = 2732

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Score = 637 bits (1644), Expect = 0.0
  Identities = 287/481 (59%), Positives = 366/481 (76%), Gaps = 5/481 (1%)
 Ouerv: 6
            MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65
            +TY RLIS+MGFK++ ++GY +FITR+EAIR VRAW+GFD EG HATRD++GTN PLO
 Sbict: 1586 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIRRVRAWVGFDAEGAHATRDSIGTNFPLO 1645
 Ouerv: 66
            LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125
            LGFSTG++ V TG + F + A+ PPG+OFKHL+PLM +G W+VVRI+IVO
 Sbjct: 1646 LGFSTGIDFVVEATGMFAERDGYVFKKAVARAPPGEQFKHLVPLMSRGQKWDVVRIRIVO 1705
Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185
            MLSD L L+D VV V WA FELT ++YF K+G E C +C+KRATCF++ + Y CW
Sbict: 1706 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGKEVVCSVCNKRATCFNSRTGYYGCWR 1765
Query: 186 HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245
                DY+YNP ++D+QQWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K
Sbict: 1766 HSYSCDYLYNPLIVDIOQWGYTGSLTSNHDLICSVHKGAHVASSDAIMTRCLAVHDCFCK 1825
Ouery: 246 RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW 305
             V+WS+EYPII +E+ VN++CR +Q ++ ++A+L +++ V +DIGNPK + CV
Sbjct: 1826 SVNWSLEYPIISNEVSVNTSCRLLQRVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1883
Ouery: 306 KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365
            KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+
Sbjct: 1884 KFYDASPV---VKSVKQFVYKYRAHKDQFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 1940
Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425
            LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC
Sbict: 1941 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 2000
Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLOS 485
            +SATCITRCNLGGAVC HA +YR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS
Sbjct: 2001 RSATCITRCNLGGAVCLKHAEDYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLQS 2060
Query: 486 L 486
Sbjct: 2061 L 2061
>gi|6625761|gb|AAF19384.1|AF201929_2 RNA-directed RNA polymerase [murine
hepatitis virus strain 2]
 gi|7739595|gb|AAF68920.1|AF207902_2 RNA-directed RNA polymerase [murine
hepatitis virus strain ML-11]
         Length = 2733
 Score = 637 bits (1643), Expect = 0.0
 Identities = 287/481 (59%), Positives = 366/481 (76%), Gaps = 5/481 (1%)
           MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65
Query: 6
           +TY RLIS+MGFK++ ++GY +FITR+EAIR VRAW+GFD EG HATRD++GTN PLO
Sbjet: 1587 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIRRVRAWVGFDAEGAHATRDSIGTNFPLO 1646
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LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125

LGFSTG++ V TG + F + A+ PPG+QFKHL+PLM +G W+VVRI+IVO

Ouerv: 66

Sbict: 1647 LGFSTGIDFVVEATGMFAERDGYVFKKAVARAPPGEOFKHLVPLMSRGOKWDVVRIRIVO 1706 Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185 MLSD L L+D VV V WA FELT ++YF K+G E C +C+KRATCF++ + Y CW Sbict: 1707 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGKEVVCSVCNKRATCFNSRTGYYGCWR 1766 Ouery: 186 HSVGFDYVYNPFMIDVOOWGFTGNLOSNHDOHCOVHGNAHVASCDAIMTRCLAVHECFVK 245 DY+YNP ++D+OOWG+TG+L SNHD C VH AHVAS DATMTRCLAVH+CF K Sbict: 1767 HSYSCDYLYNPLIVDIOOWGYTGSLTSNHDLICSVHKGAHVASSDAIMTRCLAVHDCFCK 1826 Ouerv: 246 RVDWSVEYPIIGDELRVNSACRKVOHMVVKSALLADKFPVLHDIGNPKAIKCVPOAEVEW 305 V+WS+EYPII +E+ VN++CR +O ++ ++A+L +++ V +DIGNPK + CV Sbict: 1827 SVNWSLEYPIISNEVSVNTSCRLLORVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1884 Ouerv: 306 KFYDAOPCSDKAYKIEELFYSYATHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+ Sbict: 1885 KFYDASPV---VKSVKOFVYKYEAHKDOFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 1941 Ouerv: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKOLPFFYYSDSPCESHGKOVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC Sbjct: 1942 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKOVDYVPL 2001 Ouerv: 426 KSATCITRCNLGGAVCRHHANEYROYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLOS 485 +SATCITRCNLGGAVC HA +YR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS Sbjct: 2002 RSATCITRCNLGGAVCLKHAEDYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLOS 2061 Ouerv: 486 L 486 Sbjct: 2062 L 2062 >qi|2641128|qb|AAB86818.1| RNA-directed RNA polymerase [murine hepatitis virus] Length = 2733Score = 635 bits (1637), Expect = 0.0 Identities = 286/481 (59%), Positives = 364/481 (75%), Gaps = 5/481 (1%) Ouery: 6 MTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ 65 +TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HA RD++GTN PLQ Sbict: 1587 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHAIRDSIGTNFPLO 1646 Ouerv: 66 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHLIPLMYKGLPWNVVRIKIVO 125 LGFSTG++ V TG + F + A+ PPG+OFKHLIPLM +G W+VVRI+IVO Sbjct: 1647 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEOFKHLIPLMSRGOKWDVVRIRIVO 1706 Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185 MLSD L L+D VV V WA FELT ++YF K+G E C +C KRATCF++ + Y CW Sbjct: 1707 MLSDHLADLADSVVLVTWAASFELTCLRYFAKVGREVVCSVCTKRATCFNSRTGYYGCWR 1766 Ouerv: 186 HSVGFDYVYNPFMIDVOOWGFTGNLOSNHDOHCOVHGNAHVASCDAIMTRCLAVHECFVK 245 DY+YNP ++D+OOWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K Sbict: 1767 HSYSCDYLYNPLIVDIOOWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK 1826 Ouery: 246 RVDWSVEYPIIGDELRVNSACRKVOHMVVKSALLADKFPVLHDIGNPKAIKCVPOAEVEW 305 V+W++EYPII +E+ VN++CR +O ++ ++A+L +++ V +DIGNPK + CV ++

Sbict: 1827 SVNWNLEYPIISNEVSVNTSCRLLORVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1884 Ouerv: 306 KFYDAOPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+ Sbict: 1885 KFYDASPV---VKSVKOFVYKYEAHKDQFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 1941 Ouerv: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKOLPFFYYSDSPCESHGKOVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC Sbjct: 1942 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 2001 Ouerv: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQS 485 +SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS Sbict: 2002 RSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLOS 2061 Ouerv: 486 L 486 Sbjct: 2062 L 2062 >gi|4377413|emb|CAA36202.1| open reading frame 1b (AA 1-2733) [Murine hepatitis virus Length = 2733Score = 634 bits (1636), Expect = 0.0 Identities = 286/481 (59%), Positives = 364/481 (75%), Gaps = 5/481 (1%) Ouerv: 6 MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ 65 +TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HA RD++GTN PLO Sbjct: 1587 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHAIRDSIGTNFPLO 1646 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125 Query: 66 LGFSTG++ V TG + F + A+ PPG+QFKHLIPLM +G W+VVRI+IVO Sbjct: 1647 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEOFKHLIPLMSRGOKWDVVRIRIVO 1706 Ouerv: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185 MLSD L L+D VV V WA FELT ++YF K+G E C +C KRATCF++ + Y CW Sbjct: 1707 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGREVVCSVCTKRATCFNSRTGYYGCWR 1766 Ouery: 186 HSVGFDYVYNPFMIDVQOWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245 DY+YNP ++D+OOWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K Sbict: 1767 HSYSCDYLYNPLIVDIOOWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK 1826 Query: 246 RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW 305 V+W++EYPII +E+ VN++CR +Q ++ ++A+L +++ V +DIGNPK + CV Sbjct: 1827 SVNWNLEYPIISNEVSVNTSCRLLQRVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1884 Query: 306 KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+ Sbjct: 1885 KFYDASPV---VKSVKQFVYKYEAHKDQFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 1941 Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC Sbjct: 1942 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 2001 Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQS 485 +SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS

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Sbict: 2002 RSATCITRONIGGAVCI/KHAEEYREYLESYNTATTAGFTFWVYKTFDFYNI/MNTFTRIOS 2061
Ouerv: 486 L 486
Sbict: 2062 L 2062
>gi|133592|sp|P16342|RRPB CVMA5 RNA-DIRECTED RNA POLYMERASE (ORF1B)
 gi 93916 pir | S15760 genome polyprotein - murine hepatitis virus (strain
A59)
         Length = 2733
 Score = 634 \text{ bits (1636)}. Expect = 0.0
 Identities = 286/481 (59%), Positives = 364/481 (75%), Gaps = 5/481 (1%)
Ouerv: 6
           MTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65
            +TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HA RD++GTN PLQ
Sbjct: 1587 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHAIRDSIGTNFPLO 1646
Query: 66
           LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHLIPLMYKGLPWNVVRIKIVO 125
           LGFSTG++ V TG + F + A+ PPG+0FKHLIPLM +G W+VVRI+IVO
Sbjct: 1647 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEOFKHLIPLMSRGOKWDVVRIRIVO 1706
Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185
           MLSD L L+D VV V WA FELT ++YF K+G E C +C KRATCF++ + Y CW
Sbict: 1707 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGREVVCSVCTKRATCFNSRTGYYGCWR 1766
Ouery: 186 HSVGFDYVYNPFMIDVOOWGFTGNLOSNHDOHCOVHGNAHVASCDAIMTRCLAVHECFVK 245
                DY+YNP ++D+OOWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K
Sbict: 1767 HSYSCDYLYNPLIVDIOOWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK 1826
Ouery: 246 RVDWSVEYPIIGDELRVNSACRKVOHMVVKSALLADKFPVLHDIGNPKAIKCVPOAEVEW 305
             V+W++EYPII +E+ VN++CR +O ++ ++A+L +++ V +DIGNPK + CV ++
Sbjct: 1827 SVNWNLEYPIISNEVSVNTSCRLLORVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1884
Query: 306 KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365
            KEYDA P
                         +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+
Sbjct: 1885 KFYDASPV---VKSVKOFVYKYEAHKDOFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 1941
Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425
            LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC
Sbjct: 1942 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKOVDYVPL 2001
Ouery: 426 KSATCITRCNLGGAVCRHANEYROYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLOS 485
            +SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS
Sbict: 2002 RSATCTTRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLOS 2061
Query: 486 L 486
```

>gi|26008080|ref $|NP_150073.2|$  orflab polyprotein [Bovine coronavirus] Length = 7094

Score = 633 bits (1633), Expect = e-180

L Sbict: 2062 L 2062

BNSDOCID: <WO____2004092360A2_I_>

```
Identities = 284/481 (59%), Positives = 367/481 (76%). Gaps = 5/481 (1%)
 Ouerv: 6 MTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65
             +TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLO
 Sbjct: 5948 VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLO 6007
            LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125
 Query: 66
             LGFSTG++ V TG + F + AK PPG+0FKHLIPLM +G W+VVR +IVO
 Sbict: 6008 LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEOFKHLIPLMTRGORWDVVRPRIVO 6067
 Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185
            M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW
 Sbjct: 6068 MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR 6127
 Query: 186 HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245
            HSV DY+YNP ++D+QOWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF
 Sbjct: 6128 HSVTCDYLYNPLIVDIOOWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAVYDCFCN 6187
 Query: 246 RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPOAEVEW 305
             ++W+VEYPII +EL +N++CR +Q +++K+A+L +++ +DIGNPKAI CV + ++
 Sbjct: 6188 NINWNVEYPIISNELSINTSCRVLQRVMLKAAMLCNRYTLCYDIGNPKAIACV--KDFDF 6245
 Ouery: 306 KFYDAOPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365
           KFYDAQP ++ L YS+ H D F DG+C+FWNCNVD+YP NA+VCRFDTRVL+N
Sbjct: 6246 KFYDAQPI---VKSVKTLLYSFEAHKDSFKDGLCMFWNCNVDKYPPNAVVCRFDTRVLNN 6302
Ouery: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKOLPFFYYSDSPCESHGKOVVSDIDYVPL 425
           LNLPGC+GGSLYVNKHAFHT F ++AF +LK +PFFYYSD+PC
                                                                 +DYVPL
Sbjct: 6303 LNLPGCNGGSLYVNKHAFHTKPFSRAAFEHLKPMPFFYYSDTPCVYMDGMDAKQVDYVPL 6362
Ouery: 426 KSATCITRCNLGGAVCRHHANEYROYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLOS 485
          . KSATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFT+LQS
Sbjct: 6363 KSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTKLQS 6422
Ouerv: 486 L 486
Sbjct: 6423 L 6423
>gi|15077820|gb|AAK83365.1| replicase [bovine coronavirus]
          Length = 7094
 Score = 633 bits (1633), Expect = e-180
 Identities = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)
Query: 6 MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLO 65
           +TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ
Sbjct: 5948 VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ 6007
Query: 66 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125
           LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +IVQ
Sbjct: 6008 LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ 6067
Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185
           M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW
Sbjct: 6068 MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR 6127
```

Query:	186	HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK HSV DY+YNP ++D+OOWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF	245
Sbjct:	6128	${\tt HSVTCDYLYNPLIVDIQQWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAVYDCFCN}$	6187
Query:	246	RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW ++W+VEYPII +EL +N++CR +Q +++K+A+L +++ + +DIGNPKAI CV + ++	305
Sbjct:	6188	NINWNVEYPIISNELSINTSCRVLQRVMLKAAMLCNRYTLCYDIGNPKAIACVKDFDF	6245
Query:	306	KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN KFYDAQP ++ L YS+ H D F DG+C+FWNCNVD+YP NA+VCRFDTRVL+N	365
Sbjct:	6246	KFYDAQPIVKSVKTLLYSFEAHKDSFKDGLCMFWNCNVDKYPPNAVVCRFDTRVLNN	6302
Query:	366	LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL LNLPGC+GGSLYVNKHAFHT F ++AF +LK +PFFYYSD+PC +DYVPL	425
Sbjct:	6303	${\tt LNLPGCNGGSLYVNKHAFHTKPFSRAAFEHLKPMPFFYYSDTPCVYMDGMDAKQVDYVPL}$	6362
Query:	426	KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQS KSATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFT+LQS	485
Sbjct:	6363	KSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTKLQS	6422
Query:	486	L 496	
Sbjct:	6423	L 6423	
>gi 180		2 gb AAL57305.1  replicase [bovine coronavirus]	
	ь	ength = 7094	
	= 63	ength = 7094 33 bits (1633), Expect = e-180 = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)	
Identi	= 63 ities	33 bits (1633), Expect = e-180 = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%) MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ	
Identi	= 6: ities 6	33 bits (1633), Expect = e-180 = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)	65
Identi	= 63 ities 6 5948	33 bits (1633), Expect = e-180 3 bits (1633), Expect = e-180 3 bits (1634), Positives = 367/481 (76%), Gaps = 5/481 (1%) MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ +TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ	65 6007
Identi Query: Sbjct: Query:	= 63 ties 6 5948 66	33 bits (1633), Expect = e-180 .= 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%) MTYRRLISMMGFKMNYQVMGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ +TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ	65 6007 125
Identi Query: Sbjct: Query: Sbjct:	= 60 ties 6 5948 66 6008	33 bits (1633), Expect = e-180 .= 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%) MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ HTY RLISHMGFK++ ++6Y +PIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +IVQ LGFSTGJDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN	65 6007 125 6067
Identi Query: Sbjct: Query: Sbjct: Query:	= 63 ties 6 5948 66 6008	33 bits (1633), Expect = e-180 3 bits (1633), Expect = e-180 3 bits (1633), Positives = 367/481 (76%), Gaps = 5/481 (1%)  MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ +TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ  LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +TVQ LGFSTGJDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVPRRIVQ	65 6007 125 6067 185
Identi Query: Sbjct: Query: Sbjct: Query:	= 60 ties 6 5948 66 6008 126	33 bits (1633), Expect = e-180 3 bits (1633), Positives = 367/481 (76%), Gaps = 5/481 (1%)  MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ HTY RLISHMGFKH+ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFFLQ LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQPKHLIPLMYKGLPWNVVRIKIVQ LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +TVQ LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK	65 6007 125 6067 185
Identicon Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Query:	= 6008 126 6068 186	33 bits (1633), Expect = e-180  = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)  MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ  HTY RLISHMGFK++ ++6Y +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ  VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ  LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ  LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +IVQ  LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ  MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN  M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW  MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR	65 6007 125 6067 185 6127 245
Identi Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Sbjct: Sbjct:	= 65	3 bits (1633), Expect = e-180  = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)  = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)  MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ  HTY RLISHMGFK++ ++6Y +PIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ  VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ  LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKTVQ  LGFSTGH+ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +IVQ  LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQWDVVPPRTVQ  MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN  M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW  MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATANSRTGYYGCWR  HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK  HSV DY+YNP ++D+QQWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF  HSVTCDYLYNPLIVDIQQWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAV+DCFCN  RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFFVLHDIGNPKAIKCVPQAEVEW	65 6007 125 6067 185 6127 245 6187
Identi Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Query: Sbjct: Query:	= 65	33 bits (1633), Expect = e-180 3 bits (1633), Expect = e-180 3 bits (1633), Positives = 367/481 (76%), Gaps = 5/481 (1%)  MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ +TY RLIS+MGFK++ ++6Y +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ  LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +TVQ LGFSTG1PFVVEATGLFADDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN M +D L LSD VV V WA FELT ++VF K+G E +C +C KRAT +++ + Y CW MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK HSV DY+YMP ++D+QQWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF HSVTCDYLYNPLIVDIQQWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAVYDCFCN	65 6007 125 6067 185 6127 245 6187 305
Identi Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Query: Sbjct: Query:	= 6: 6: 6: 5948 66 6008 126 6068 186 6128 246 6188	3 bits (1633), Expect = e-180  = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)  = 284/481 (59%), Positives = 367/481 (76%), Gaps = 5/481 (1%)  MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ  HTY RLISHMGFK++ ++6Y +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ  VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ  LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKTVQ  LGFSTGH+ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +IVQ  LGFSTGIDFVVEATGLFADDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRTVQ  MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN  M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW  MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR  HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK  HSV DY+YNP ++D+QQWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF  HSVTCDYLYNPLIVDIQQWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAVYDCFCN  RVDWSVEYPLIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW  +WHYEYPII +EL +N++CR +Q +++K+A+L +++ + DIGNPKAI CV + ++  NINWNVEYPIISNELSINTSCRVLQRVMLKAAMLCNRYTLCYDIGNPKAIACVKDFDF	65 6007 125 6067 185 6127 245 6187 305

Ouery: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKOVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF +LK +PFFYYSD+PC +DYVPI Sbict: 6303 LNLPGCNGGSLYVNKHAPHTKPFSRAAFEHLKPMPFFYYSDTPCVYMDGMDAKOVDYVPL 6362 Ouery: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLOS 485 KSATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFT+LOS Sbjct: 6363 KSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTKLOS 6422 Query: 486 L 486 т. Sbict: 6423 T. 6423 >gi|7769353|gb|AAF69342.1|AF208067_2 RNA-directed RNA polymerase [murine hepatitis virusl Length = 2733Score = 633 bits (1633), Expect = e-180 Identities = 285/481 (59%), Positives = 364/481 (75%), Gaps = 5/481 (1%) Ouery: 6 MTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ 65 ++Y RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HA RD++GTN PLO Sbjct: 1587 VSYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHAIRDSIGTNFPLQ 1646 Ouerv: 66 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125 LGFSTG++ V TG + F + A+ PPG+QFKHLIPLM +G W+VVRI+IVO Sbjct: 1647 LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEOFKHLIPLMSRGOKWDVVRIRIVO 1706 Ouery: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185 MLSD L L+D VV V WA FELT ++YF K+G E C +C KRATCF++ + Y CW Sbjct: 1707 MLSDHLVDLADSVVLVTWAASFELTCLRYFAKVGREVVCSVCTKRATCFNSRTGYYGCWR 1766 Query: 186 HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245 DY+YNP ++D+QQWG+TG+L SNHD C VH AHVAS DAIMTRCLAVH+CF K Sbjct: 1767 HSYSCDYLYNPLIVDIQQWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK 1826 Query: 246 RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW 305 V+W++EYPII +E+ VN++CR +Q ++ ++A+L +++ V +DIGNPK + CV Sbjct: 1827 SVNWNLEYPIISNEVSVNTSCRLLQRVMFRAAMLCNRYDVCYDIGNPKGLACVKG--YDF 1884 Query: 306 KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDA P +++ Y Y H D+F DG+C+FWNCNVD+YPANA+VCRFDTRVL+ Sbjct: 1885 KFYDASPV---VKSVKOFVYKYEAHKDOFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK 1941 Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425 LNLPGC+GGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC Sbjct: 1942 LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL 2001 Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQS 485 +SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLOS Sbjct: 2002 RSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLQS 2061

Query: 486 L 486

__2004092360A2_l_s

Sbjct: 2062 L 2062

Score = 623 bits (1607), Expect = e-177Identities = 282/481 (58%), Positives = 365/481 (75%), Gaps = 5/481 (1%)

Query: 6 MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ 65 +TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLO

Sbjct: 1574 VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ 1633

Query: 66 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125 LGFSTG++ V TG + F + AK PPG+OFKHLIPLM +G W+VVR +IVO

Sbjct: 1634 LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ 1693

Query: 126 MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN 185

M +D L LSD VV V WA FELT ++YF K+G E +C + KRAT +++ + Y CW Sbict: 1694 MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVSTKRATAYNSRTGYYGCWR 1753

Query: 186 HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK 245 HSV DY+YNP ++D+QQWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF

Sbjct: 1754 HSVTCDYLYNPLIVDIQQWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAVYDCFCN 1813

Query: 246 RVDWSVEYPTIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW 305 . ++W+VEYPII +EL +N++CR +Q +++K+A+L +++ + +DIGNPKAI CV +++

Sbjct: 1814 NINWNVEYPIISNELSINTSCRVLORVMLKAAMLCNRYTLCYDIGNPKAIACV--KDFDF 1871

Query: 306 KFYDAQPCSDKAYKIRELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN 365 KFYDAQP ++ L Y + H D F DG+C+FWNCNVD+YP NA+VCRFDTRVL+N

Sbjct: 1872 KFYDAQPI---VKSVKTLLYFFEAHKDSFKDGLCMFWNCNVDKYPPNAVVCRFDTRVLNN 1928

Query: 366 "LNLPGCDGGSLYUNKHAPHTPAFPKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425 "
BALFGC+GGSLYVNKHAPHT F ++AF +LK +PFFYYSD+PC +DYVPL Sbict: 1929, LNLPGCHGGSLYVNKHAPHTKPFSRAAFEHLKFMPFFYYSDTPCVYMDGMDAKOVDYVPL 1988

Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQS 485 KSATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFT+LQS

Sbjct: 1989 KSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTKLQS 2048

Query: 486 L 486

Sbjct: 2049 L 2049

>gi|25121571|ref|NP_740618.1| coronavirus nsp11 [Murine hepatitis virus]
Length = 521

Score = 622 bits (1603), Expect = e-177
Identities = 284/479 (59%), Positives = 362/479 (75%), Gaps = 5/479 (1%)

Query: 6 MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ 65

+TY RLIS+MGFK++ ++GY +FITR+EAI+ VRAW+GFD EG HA RD++GTN PLQ
Sbjct: 48 VTYSRLISLMGFKLDLTLDGYCKLFITRDEAIKRVRAWVGFDAEGAHAIRDSIGTNFPLQ 107

Query: 66 LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQ 125

BNSDOCID: <WO_

_2004092360A2_l_>

# 167/193

٤	Bbjct:	108	LGFSTGIDFVVEATGMFAERDGYVFKKAAARAPPGEQFKHLIPLMSRGQKWDVVRIRIVQ	167
Ç	uery:	126	MLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWN	185
5	bjct:	168	MLSD L L+D VV V WA FELT ++YF K+G E C +C KRATCF++ + Y CW MLSDHLADLADSVVLVTWAASFELTCLRYFAKVGREVVCSVCTKRATCFNSRTGYYGCWR	227
Ç	uery:	186	HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK HS DY+YNP ++D+OOWG+TG+I, SNHD, C, VH, ANNAC, DAIMTRCLAVHECFVK	245
			HSYSCDYLYNPLIVDIQQWGYTGSLTSNHDPICSVHKGAHVASSDAIMTRCLAVHDCFCK	287
Ç	uery:	246	RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW V+W++EYPII +E+ VN++CR +Q ++ ++A+L +++ V +DIGNPK + CV ++	305
			SVNWNLEYPIISNEVSVNTSCRLLQRVMFRAAMLCNRYDVCYDIGNPKGLACVKGYDF	
Ç	uery:	306	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	365
			KFYDASPVVKSVKQFVYKYEAHKDQFLDGLCMFWNCNVDKYPANAVVCRFDTRVLNK	
Ç	uery:	366	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	425
S	bjct:	403	LNLPGCNGGSLYVNKHAFHT F ++AF NLK +PFFYYSD+PC +DYVPL LNLPGCNGGSLYVNKHAFHTSPFTRAAFENLKPMPFFYYSDTPCVYMEGMESKQVDYVPL	462
Q	uery:	426	KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQ	484
	•		+SATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFTRLQ RSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTRLQ	
			92 ref NP_742140.1  coronavirus nsp11 [Bovine coronavirus] Length = 521	
	Score Identi	= 6	617 bits (1590), Expect = e-175 s = 282/479 (58%), Positives = 365/479 (76%), Gaps = 5/479 (19	<b>b</b> )
Q	uery:	6	MTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQ	65
S	bjct:	48	+TY RLIS+MGFK++ ++GY +FIT+EEA++ VRAW+GFD EG HATRD++GTN PLQ VTYSRLISLMGFKLDVTLDGYCKLFITKEEAVKRVRAWVGFDAEGAHATRDSIGTNFPLQ	107
Q	uery:	66	LGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKĮVQ	125
			LGFSTG++ V TG + F + AK PPG+QFKHLIPLM +G W+VVR +LVQ LGFSTGIDFVVEATGLFADRDGYSFKKAVAKAPPGEQFKHLIPLMTRGQRWDVVRPRIVQ	
		126	MLSDTLKGLSDRVVFVLWAHGFELTSMKVFVKTGDERTCCLCDKDATCECTGCDTVACTAN	
S	bjct:	168	M +D L LSD VV V WA FELT ++YF K+G E +C +C KRAT +++ + Y CW MFADHLIDLSDCVVLVTWAANFELTCLRYFAKVGREISCNVCTKRATAYNSRTGYYGCWR	227
Q۱	uery:	186	HSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVK	245
			HSV DY+YNP ++D+QQWG+ G+L SNHD +C VH AHVAS DAIMTRCLAV++CF HSVTCDYLYNFLIVDIQQWGYIGSLSSNHDLYCSVHKGAHVASSDAIMTRCLAVYDCFCN	
Q۱	ery:	246	RVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEW	305
			++W+VEYPII +EL +N++CR +Q +++K+A+L +++ + +DIGNPKAI CV + ++ NINWNVEYPIISNELSINTSCRVLQRVMLKAAMLCNRYTLCYDIGNPKAIACVKDFDF	
		306	KFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLSN KFYDAQP ++ L YS+ H D F DG+C+FWNCNVD+YP NA+VCRFDTRVL+N	

Sbjct: 346 KFYDAQPIVKSVKTLLYSFEAHKDSFKDGLCMFWNCNVDKYPPNAVVCRFDTRVLNN 402	
Query: 366 LNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVPL 425 LNLPGC+GGŠLYVNKHAFHT F ++AF +LK +PFFYYSD+PC +PYVPL	
Sbjct: 403 LNLPGCNGGSLYVNKHAFHTKPFSRAAFEHLKPMPFFYYSDTPCVYMDGMDAKQVDYVPL 462	
Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQ 484 KSATCITRCNLGGAVC HA EYR+YL++YN +AGF+ W+YK FD YNLWNTFT+LO	
Sbjct: 463 KSATCITRCNLGGAVCLKHAEEYREYLESYNTATTAGFTFWVYKTFDFYNLWNTFTKLQ 521	
>gi   10242469   ref   NP_066134.1   ORFlab polyprotein; frameshift product [Avi- infectious bronchitis virus] Length = 6629	an
Score = 575 bits (1482), Expect = e-163	
Identities = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)	
Query: 5 DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL 64 ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA +GTNLP	
Sbjct: 5515 EİTYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLPF 55	74
Query: 65 QLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIV 12 Q+GFSTG + V P G VDT F VN+K PPG+QF HL L PW+V+R +IV	4
Sbjct: 5575 QVGFSTGADFVVTPEGLVDTSIGNNFEPVNSKAPPGEQFNHLRVLFKSAKPWHVIRPRIV 56	34
Query: 125 QMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACW 18 QML+D L +SD VVFV W HG ELT+++YFVKIG E+ C C RAT F++ + YACW	4
Sbjct: 5635 QMLADNLCNVSDCVVFVTWCHGLELTTLRYFVKIGKEQVCS-CGSRATTFNSHTQAYACW 56	93
Query: 185 NHSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFV 24 H +GFD+VYNP ++D+QQWG++GNLQ NHD HC VHG+AHVAS DAIMTRCLA++ F	4
Sbjct: 5694 KHCLGFDFVYNPLLVDIQQWGYSGNLQFNHDLHCNVHGHAHVASVDAIMTRCLAINNAFC 57	53
Query: 245 KRVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVB 30 + V+W + YP I +E VNS+CR +Q M + + + A K V++DIGNPK IKCV + +V	4
Sbjct: 5754 QDVNWDLTYPHIANEDEVNSSCRYLQRMYLNACVDALKVNVVYDIGNPKGIKCVRRGDVN 58	13
Query: 305 WKFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLS 36 ++FYD P + E Y Y H DKF DG+C+FWNCNVD YP N++VCR+DTR LS	4
Sbjct: 5814 FRFYDKNPIVRNVKQFEYDYNQHKDKFADGLCMFWNCNVDCYPDNSLVCRYDTRNLS 58	70
Query: 365 NLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVP 42	4
NLPGC+GGSLYVNKHAF+TF FD+ +F NLK +PFF+Y SPCE+ V+ D V Sbjct: 5871 VFNLPGCNGGSLYVNKHAFYTPKFDRISFRNLKAMPFFFYDSSPCETIQVDGVAQ-DLVS 59	29
Query: 425 LKSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQ 48	4
L + CIT+CN+GGAVC+ HA Y +++ +YN ++AGF+ W+ + + YNLW +F+ LQ Sbjct: 5930 LATKDCITKCNIGGAVCKKHAQMYAEFVTSYNAAVTAGFTFWVINKLNPYNLWKSFSALQ 59	89
Query: 485 SL 486	
Shict: 5990 SI 5991	

>gi|14149033|emb|CAC39112.1| replicase polyprotein lab [Avian infectious bronchitis virus (strain

Beaudette CK)]
Lenoth = 6629

Score = 575 bits (1482), Expect = e-163 Identities = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)

Query: 5 DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL 64 ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA +GTNLP

Sbjct: 5515 EITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLPF 5574

Query: 65 QLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIV 124 Q+GFSTG + V P G VDT F VN+K PPG+0F HL L PW+V+R +TV

Sbjct: 5575 QVGFSTGADFVVTPEGLVDTSIGNNFEPVNSKAPPGEQFNHLRVLFKSAKPWHVIRPRIV 5634

Query: 125 QMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACW 184 QML+D L +SD VVFV W HG ELT+++YFVKIG E+ C C RAT F++ + YACW

Sbjct: 5635 QMLADNLCNVSDCVVFVTWCHGLELTTLRYFVKIGKEQVCS-CGSRATTFNSHTQAYACW 5693

Query: 185 NHSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFV 244 H +GFD+VYNP ++D+QQWG++GNLQ NHD HC VHG+AHVAS DAIMTRCLIA++ F

Sbjct: 5694 KHCLGFDFVYNPLLVDIQQWGYSGNLQFNHDLHCNVHGHAHVASVDAIMTRCLAINNAFC 5753

Query: 245 KRVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVE 304

+ V+W + YP I +E VNS+CR +Q M + + + A K V++DIGNPK IKCV + +V
Sbjct: 5754 QDVNWDLTYPHIANEDEVNSSCRYLQRMYLNACVDALKVNVVYDIGNPKGIKCVRRGDVN 5813

Query: 305 WKFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLS 364

++FYD P + E Y Y H DKF DG+C+FWNCNVD YP N++VGR+DTR LS
Sbjct: 5814 FRFYDKNPIVRNVKQFE---YDYNQHKDKFADGLCMFWNCNVDCYPDNSLVCRYDTRNLS 5870

Query: 365 NLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVP 424 NLPGC+GGSLYVNKHAF+TP FD+ +F NLK +PFF+Y SPCE+ V+ D V

NDEGC+GGSLYVNKHAF+TP FD+ +F NLK +PFF+Y SPCE+ V+ D V
Sbjct: 5871 VFNLPGCNGGSLYVNKHAFYTPKFDRISFRNLKAMPFFFYDSSPCETIQVDGVAQ-DLVS 5929

Query: 425 LKSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQ 484 L + CIT+CN+GGAVC+ HA Y ++++YN ++AGF+ W+ + YNLW +F+ LO

Sbjct: 5930 LATKDCITKCNIGGAVCKKHAQMYAEFVTSYNAAVTAGFTFWVTNKLNPYNLWKSFSALQ 5989

Query: 485 SL 486

Sbjct: 5990 SI 5991

>gi|458735|emb|CAA83018.1| potential chimeric protein [Avian infectious bronchitis virus] Length = 2155

Score = 570 bits (1470), Expect = e-161 Identities = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)

Query: 5 DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL 64 ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA +GTNLP Sbjct: 1596 EITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLPF 1655

Query:		QLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIV Q+GFSTG + V P G VDT F VN+K PPG+QF HL L PW+V+R +IV	
Sbjct:	1656	${\tt QVGFSTGADFVVTPEGLVDTSIGNNFEPVNSKAPPGEQFNHLRVLFKSAKPWHVIRPRIV}$	1715
Query:	125	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	184
Sbjct:	1716	QMLADNLCNVSDCVVFVTWCHGLELTTLRYFVKIGKEQVCS-CGSRATTFNSHTQAYACW	1774
Query:	185	NHSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFV H +GFD+VYNP ++D+QQWG++GNLQ NHD HC VHG+AHVAS DAIMTRCLA++ F	244
Sbjct:	1775	KHCLGFDFVYNPLLVDIQQWGYSGNLQFNHDLHCNVHGHAHVASVDAIMTRCLAINNAFC	1834
Query:	245	KRVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVE + V+W + YP I +E	304
Sbjct:	1835	QDVNWDLTYPHIANEDEVNSSCRYLQRMYLNACVDALKVNVVYDIGNPKGIKCVRRGDVN	1894
Query:	305	WKFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLS ++FYD P + E Y Y H DKF DG+C+FWNCNVD YP N++VCF+DTR LS	364
Sbjct:	1895	++FYD P + E Y Y H DKF DG+C+FWNCNVD YP N++VCR+DTR LS FRFYDKNPIVRNVKQFEYDYNQHKDKFADGLCMFWNCNVDCYPDNSLVCRYDTRNLS	1951
Query:	365	NLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSDIDYVP NLPGC+GGSLYVNKHAF+TP FD+ +F NLK +PFF+Y SPCE+ V+ D V	424
Sbjct:	1952	NIPGC+GGSLYVNKHAF+TP FD+ +F NLK +PFF+Y SPCE+ V+ D V VFNLPGCNGGSLYVNKHAFYTPKFDRISFRNLKAMPFFFYDSSPCETIQVDGVAQ-DLVS	2010
Query:	425	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	484
Sbjct:	2011	LATKOCITKCNIGGAVCKKHAQMYAEFVTSYNAAVTAGFTFWVTNKLNPYNLWKSFSALQ	2070
Query:	485		
		SL 486 S+ SI 2072	
Sbjct: >gi   133 gi   748 virus gi   292	2071 3594 s 326 p3 (stra	S+ SI 2072  SP P26314 RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORF1E) LT  VF1HB2 genome polyprotein - avian infectious bronchitis	
Sbjct: >gi   133 gi   748 virus gi   292 gi   333	2071 3594 s 326 p3 (stra 2953 s 1173 s Le	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORF1B) LT   VFIHB2 genome polyprotein - avian infectious bronchitis ain  Beaudette) JB   AAA70234 1   pol protein [Avian infectious bronchitis virus] JB   AAA46224.1   ORF1b [Avian infectious bronchitis virus]	
Sbjct: >gi   133 gi   748 virus gi   292 gi   333	2071 3594 s 326 p (stra 2953 s 1173 s Le = 57	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORF1B) ir     VFIHB2 genome polyprotein - avian infectious bronchitis ain  Beaudette) jb   AAA70234.1   pol protein [Avian infectious bronchitis virus] be   AAA46224.1   ORF1b [Avian infectious bronchitis virus] bength = 2652  70 bits (1469), Expect = e-161 = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)  DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL	
Sbjct:  >gi   133 gi   746 virus  gi   292 gi   333  Score Identi Query:	2071 3594 s 326 p3 (stra 2953 s 1173 s Le = 53	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORFIB) II   VFIHB2 genome polyprotein - avian infectious bronchitis ain Beaudette) Jb   AAA70234.1   pol protein [Avian infectious bronchitis virus] Beaha46224.1   ORFIb [Avian infectious bronchitis virus] Bength = 2652  Obits (1469), Expect = e-161 = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)	64
Sbjct:  >gi   133 gi   746 virus  gi   292 gi   333  Score Identi Query:	2071 3594   5 326   po (stra 2953   c 1173   c Le = 57 ities 5 1538	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA FOLYMERASE (ORF1E) LT     VFIHB2 genome polyprotein - avian infectious bronchitis ain Beaudette) Beaudette) Beaudette) Beaudette   Double	64 1597
Sbjct:  >gi   133 gi   748 virus gi   292 gi   333  Score Identi Query: Sbjct: Query:	2071 3594 s 326 p (stra 2953 s 173 s Le = 55 ities 5 1538	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORF1B) ir     VFIHB2 genome polyprotein - avian infectious bronchitis ain  Beaudette) jb   AAA70234.1   pol protein [Avian infectious bronchitis virus] ength = 2652  70 bits (1469), Expect = e-161 = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)  DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA +GTNLP EITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNLGTNLPF	64 1597 124
Sbjct:  >gi   133 gi   748 virus gi   292 gi   333  Score Identi Query: Sbjct: Query:	2071 8594   8 326   pa (stra 2953   9 1173   9 1 ties 5 1538 65 1598	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORF1B) ix     VFIHB2 genome polyprotein - avian infectious bronchitis ain  Beaudette) jb   AAA70234.1   pol protein [Avian infectious bronchitis virus] ength = 2652  70 bits (1469), Expect = e-161 = 262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)  DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA +GTNLP EITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLFF QLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPFGDQFKHLIPLMYKGLFMNVVRIKIV QVGFSTGADFVVTPEGLVDTSIGNNFEPVNSKAPPGEQFNHLRVLFKSAKPWHVIRPRIV QMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACW	64 1597 124 1657
Sbjct:  ygi   133 gi   746 virus  gi   292 gi   333  Score Identi Query: Sbjct: Query: Sbjct: Query:	2071 3594 s 326 p (stra (stra 1578 s 1538 65 1598 125	S+ SI 2072  SP   P26314   RRPB_IBVB RNA-DIRECTED RNA POLYMERASE (ORF1B) LT     VFIHB2 genome polyprotein - avian infectious bronchitis ain Beaudette) Beaudette) Beaudette) Beaudette) Beaudette) Bo   AAA70234.1   pol protein [avian infectious bronchitis virus] Beaudette) Bo   AAA70234.1   pol protein [avian infectious bronchitis virus] Bength = 2652  Obits (1469), Expect = e-161   262/482 (54%), Positives = 344/482 (71%), Gaps = 5/482 (1%)  DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL H+TT+ LISH+GFKM+ V G NMFITR+EAIRHVRAWIGFDVE HA +GTNLPL BITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLPF CUGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLFWNVVRIKIV Q+GFSTG+ V P G VDT F VN+K PPG+QF HL L PW+V+R +IV QVGFSTGADFVVTPEGLVDTSIGNNFEPVNSKAPPGEQFNHLRVLFKSAKFWHVIRPRIV	64 1597 124 1657

BNSDOCID: <WO____2004092360A2_I_>

Query: 185 NHSVGFDYVYNPFMIDVQOWGFTGNLOSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFV 244 H +GFD+VYNP ++D+QQWG++GNLQ NHD HC VHG+AHVAS DAIMTRCLA++ F Sbjct: 1717 KHCLGFDFVYNPLLVDIQOWGYSGNLQFNHDLHCNVHGHAHVASVDAIMTRCLAINNAFC 1776 Ouery: 245 KRVDWSVEYPIIGDELRVNSACRKVOHMVVKSALLADKFPVLHDIGNPKAIKCVPOAEVE 304 + V+W + YP I +E VNS+CR +O M + + + A K V++DIGNPK IKCV + +V Sbjct: 1777 QDVNWDLTYPHIANEDEVNSSCRYLQRMYLNACVDALKVNVVYDIGNPKGIKCVRRGDVN 1836 Query: 305 WKFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLS 364 ++FYD P + E Y Y H DKF DG+C+FWNCNVD YP N++VCR+DTR LS Sbjct: 1837 FRFYDKNPIVRNVKQFE---YDYNQHKDKFADGLCMFWNCNVDCYPDNSLVCRYDTRNLS 1893 Query: 365 NLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKOLPFFYYSDSPCESHGKOVVSDIDYVP 424 NLPGC+GGSLYVNKHAF+TP FD+ +F NLK +PFF+Y SPCE+ Sbjct: 1894 VFNLPGCNGGSLYVNKHAFYTPKFDRISFRNLKAMPFFFYDSSPCETIQVDGVAQ-DLVS 1952 Query: 425 LKSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLO 484 L + CIT+CN+GGAVC+ HA Y +++ +YN ++AGF+ W+ + + YNLW +F+ LO Sbjet: 1953 LATKDCITKCNIGGAVCKKHAOMYAEFVTSYNAAVTAGFTFWVTNKLNPYNLWKSFSALO 2012 Ouery: 485 St 486 ... Sbjct: 2013 SI 2014 >gi|29293454|gb|AA067706.1| ORF1b polyprotein [Avian infectious bronchitis virusl Length = 2649 Score = 565 bits (1455), Expect = e-160 Identities = 261/482 (54%), Positives = 342/482 (70%), Gaps = 8/482 (1%) Ouerv: 5 DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL 64 ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA Sbjct: 1538 EITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLPF 1597 Ouerv: 65 QLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIV 124 Q+GFSTG + V P G +DT F VN+K PPG+OF HL L PW+V+R +IV Sbjct: 1598 QVGFSTGADFVVTPEGLIDTSIGNNFEPVNSKAPPGEQFNHLRALFKSAKPWHVIRPRIV 1657 Query: 125 QMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACW 184 OML+D L +SD VVFV W HG ELT+++YFVKIG E+ C C RAT F++ + YACW Sbjct: 1658 QMLADNLCNVSDCVVFVTWCHGLELTTLRYFVKIGKEQVCS-CGSRATTFNSHTOAYACW 1716 Query: 185 NHSVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFV 244 VYNP ++D+QQWG++GNLQ NHD HC VHG+AHVAS DA+MTRCLA++ F Sbjct: 1717 RHCLG---VYNPLLVDIQQWGYSGNLQFNHDLHCNVHGHAHVASADAVMTRCLAINNAFC 1773 Query: 245 KRVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVE 304 K V+W ++YP I +E VNS+CR +Q M + + + A K V++DIGNPK IKCV + +V Sbjct: 1774 KDVNWELQYPHIANEDEVNSSCRYLQRMYLNACVDALKVNVVYDIGNPKGIKCVRRGDVN 1833 Query: 305 WKFYDAQFCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLS 364 + E Y Y+ H DKF DG+C+FWNCNVD YP N++VCR+DTR LS Sbjct: 1834 FRFYDKNPIVPNVKQFE---YDYSQHKDKFADGLCMFWNCNVDCYPENSLVCRYDTRNLS 1890

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Query: 365 NLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKOVVSDIDYVP 424
             NLPGC+GGSLYVNKHAFHTP FD+ +F NLK +PFF+Y SPCE+
Sbict: 1891 VFNLPGCNGGSLYVNKHAFHTPKFDRISFRNLKAMPFFFYDSSPCETIOVDGVAO-DLVS 1949
Ouery: 425 LKSATCITRCNLGGAVCRHHANEYROYLDAYNMMISAGFSLWIYKOFDTYNLWNTFTRLO 484
           L + CIT+CN+GGAVC+ HA Y +++ +YN ++AGF+ W+
                                                         F+ YNLW F+ LO
Sbict: 1950 LATKDCITKCNIGGAVCKKHAOMYAEFVFSYNAAVTAGFTFWVTNNFNPYNLWKNFSALQ 2009
Ouerv: 485 SL 486
           G.
Sbict: 2010 SI 2011
>qi|25121555|ref|NP_740631.1| coronavirus nsp11 [Avian infectious
bronchitis virus
          Length = 521
 Score = 559 bits (1440), Expect = e-158
 Identities = 261/480 (54%), Positives = 342/480 (71%), Gaps = 5/480 (1%)
Ouerv: 5
          DMTYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPL 64
           ++TY+ LIS++GFKM+ V G NMFITR+EAIR+VR W+GFDVE HA
                                                               +CTIMIT.D
Sbict: 47 EITYKHLISLLGFKMSVNVEGCHNMFITRDEAIRNVRGWVGFDVEATHACGTNIGTNLPF 106
Ouery: 65 OLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHLIPLMYKGLPWNVVRIKIV 124
           O+GFSTG + V P G VDT
                                   F VN+K PPG+OF HL L
                                                            PW+V+R +TV
Sbjct: 107 QVGFSTGADFVVTPEGLVDTSIGNNFEPVNSKAPPGEQFNHLRVLFKSAKPWHVIRPRIV 166
Query: 125 QMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACW 184
           OML+D L +SD VVFV W HG ELT+++YFVKIG E+ C C RAT F++ + YACW
Sbjct: 167 QMLADNLCNVSDCVVFVTWCHGLELTTLRYFVKIGKEQVCS-CGSRATTFNSHTQAYACW 225
Ouery: 185 NHSVGFDYVYNPFMIDVOOWGFTGNLOSNHDOHCOVHGNAHVASCDAIMTRCLAVHECFV 244
           H +GFD+VYNP ++D+OOWG++GNLQ NHD HC VHG+AHVAS DAIMTRCLA++ F
Sbjct: 226 KHCLGFDFVYNPLLVDIOOWGYSGNLOFNHDLHCNVHGHAHVASVDAIMTRCLAINNAFC 285
Ouerv: 245 KRVDWSVEYPIIGDELRVNSACRKVOHMVVKSALLADKFPVLHDIGNPKATKCVPOAEVE 304
           + V+W + YP I +E VNS+CR +Q M + + + A K V++DIGNPK IKCV + +V
Sbjct: 286 QDVNWDLTYPHIANEDEVNSSCRYLORMYLNACVDALKVNVVYDIGNPKGIKCVRRGDVN 345
Ouery: 305 WKFYDAOPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRVLS 364,
                         + E
                              Y Y H DKF DG+C+FWNCNVD YP N++VCR+DTR LS
Sbjct: 346 FRFYDKNPIVRNVKQFE---YDYNQHKDKFADGLCMFWNCNVDCYPDNSLVCRYDTRNLS 402
Query: 365 NLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKOLPFFYYSDSPCESHGKOVVSDIDYVP 424
             NLPGC+GGSLYVNKHAF+TP FD+ +F NLK +PFF+Y SPCE+
Sbjct: 403 VFNLPGCNGGSLYVNKHAFYTPKFDRISFRNLKAMPFFFYDSSPCETIOVDGVAO-DLVS 461
Query: 425 LKSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRLQ 484
           L + CIT+CN+GGAVC+ HA Y +++ +YN ++AGF+ W+ + + YNLW +F+ LQ
Sbict: 462 LATKDCITKCNIGGAVCKKHAOMYAEFVTSYNAAVTAGFTFWVTNKLNPYNLWKSFSALO 521
>gi|9635157|ref|NP_058422.1| replicase [Transmissible gastroenteritis
virus]
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gi|7801348|emb|CAB91143.1| replicase [Transmissible gastroenteritis virus]

Length = 6685

Score = 545 bits (1403), Expect = e-153 Identities = 261/484 (53%), Positives = 335/484 (69%), Gaps = 13/484 (2%) Ouerv: 4 KDMTYRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLP 63 KD+ Y +IS MGF+ + GY +F TR+ A+R+VRAW+GFDVEG H Sbjct: 5574 KDVKYANVISYMGFRFEANIPGYHTLFCTRDFAMRNVRAWLGFDVEGAHVCGDNVGTNVP 5633 LQLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKI 123 Ouerv: 64 LOLGES GV+ V GVTE V A+ PPG+OF HLIPLM KG PW++VR +I Sbjct: 5634 LQLGFSNGVDFVVQTEGCVITEKGNSIEVVKARAPPGEQFAHLIPLMRKGOPWHIVRRRI 5693 Query: 124 VQMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIG-PERTCCLCDKRATCFSTSSDTYA 182 GLSD ++FVLWA G ELT+M+YFVKIG P++ C C K ATC+S+S Sbjct: 5694 VQMVCDYFDGLSDILIFVLWAGGLELTTMRYFVKIGRPQK--CECGKSATCYSSSOSVYA 5751 Ouery: 183 CWNHSVGFDYVYNPFMIDVOOWGFTGNLOSNHDOHCOVHGNAHVASCDAIMTRCLAVHEC 242 C+ H++G DY+YNP+ ID+OOWG+TG+L NH + C +H N HVAS DAIMTRCLA+H+C Sbjct: 5752 CFKHALGCDYLYNPYCIDIQOWGYTGSLSMNHHEVCNIHRNEHVASGDAIMTRCLAIHDC 5811 Query: 243 FVKRVDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAE 302 FVKRVDWS+ YP I +E ++N A R VQ V+K+AL +HD+GNPK I+C Sbjct: 5812 FVKRVDWSIVYPFIDNEEKINKAGRIVOSHVMKAALKIFNPAAIHDVGNPKGIRCA-TTP 5870 Ouery: 303 VEWKFYDAQPCSDKAYKIEELFYSYAIHHDKFTDGVCLFWNCNVDRYPANAIVCRFDTRV 362 + W YD P ++ + L Y Y +H +G+ LFWNCNVD YP +IVCRFDTR Sbjct: 5871 IPWFCYDRDPINN---NVRCLDYDYMVHGQ--MNGLMLFWNCNVDMYPEFSIVCRFDTRT 5925 Ouery: 363 LSNLNLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKOLPFFYYSDSPCESHGKOVVSDIDY 422 S L+L GC+GG+LYVN HAFHTPA+D+ AF LK +PFFYY DS CE Sbjct: 5926 RSKLSLEGCNGGALYVNNHAFHTPAYDRRAFAKLKPMPFFYYDDSNCE----LVDGQPNY 5981 Ouery: 423 VPLKSATCITRCNLGGAVCRHHANEYROYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTR 482 VPLKS CIT+CN+GGAVC+ HA YR Y++ YN+ + AGF++W + FDTY LW+ F Sbjct: 5982 VPLKSNVCITKCNIGGAVCKKHAALYRAYVEDYNIFMQAGFTIWCPQNFDTYMLWHGFVN 6041 Query: 483 LOSL 486

++L Sbjct: 6042 SKAL 6045

>gi | 19387582 | ref|NP_598309.1 | Foll [porcine epidemic diarrhea virus]
gi | 13752450 | gb | AAK38661.1 | Foll [porcine epidemic diarrhea virus]
Length = 6781

Score = 541 bits (1394), Expect = e-152 Identities = 256/480 (53%), Positives = 334/480 (69%), Gaps = 12/480 (2%)

Query: 8 YRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQLG 67 Y +IS MGF+ + + + + F TR+ A+R+VR W+GFDVEG H VGTN+PLQLG

Sbjct: 5675 YEHVISFMGFRFDINIPNHHTLFCTRDFAMRNVRGWLGFDVEGAHVVGSNVGTNVPLQLG 5734
Query: 68 FSTGVNLVAVPTGYVLVTFMNMFERMUNININDSSOCRETARION FOR FAMILY AND PROPERTY AND

Query: 68 FSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQML 127
FS GV+ V P G V TE+ V A+ PPG+QF HL+PL+ +G PW+VVR +IVQM
Sbjct: 5735 FSNGVDFVVRPEGCVVTESGDYIKPVRARAPPGEQFAHLLPLLKRGQPWDVVRKRIVQMC 5794

# 174/193 Ouery: 128 SDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWNHS 187

	Sbjct:		SD L LSD ++FVLWA G ELIT+M+YFVKIGP ++C C K ATC++++ TY C+ H+ SDYLANLSDILIFVLWAGGLELTTMRYFVKIGPSKSCD-CGKVATCYNSALHTYCCFKHA	5853
	Query:	188	VGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVKRV	247
	Sbjct:	5854	+G DY+YNP+ ID+QQWG+ G+L NH +HC VH N HVAS DAIMTRCLA+H+CFVK V LGCDYLYNPYCIDIQQWGYKGSLSLNHHEHCNVHRNEHVASGDAIMTRCLAIHDCFVKNV	5913
	Query:		DWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFFVLHDIGNPKAIKCVPQAEVEWKF DWS+ YP IG+E +N + R VO ++S L ++DIGNPK I+C + +W	307
	Sbjct:		DWSITYPFIGNEAVINKSGRIVQSHTMRSVLKLYNPKAIYDIGNPKGIRCA-VTDAKWFC	5972
	Query:	308	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	367
	Sbjct:	5973	FDKNPTNSNVKTLEYDY-ITHGQF-DGLCLFWNCNVDMYPEFSVVCRFDTRCRSPLN	6027
	Query:	368	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	427
	Sbjct:	6028	${\tt LEGCNGGSLYVNNHAFHTPAFDKRAFAKLKPMPFFFYDDTECDKLQDSINYVPLRA}$	6083
			$ \begin{array}{llllllllllllllllllllllllllllllllllll$	
	Sbjct:	6084	${\tt SNCITKCNVGGAVCSKHCAMYHSYVNAYNTFTSAGFTIWVPTSFDTYNLWQTFSNNLQGL}$	6143
	229E1		7   ref   NP_073549.1   replicase polyprotein lab [Human coronavi	
	corona	virus L	0 gb AAG48591.1 AF304460_2 replicase polyprotein lab [Human 229E] : ength = 6758	
	corona	virus Le = 5	229E]	
	Score Ident	virus Lo = 5: ities 7	229B; ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (23  TYRRLISMMGFKMNYQVNGYFNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY +1S MGF+ + + G +++ TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+	s) 66
	Score Ident. Query: Sbjct:	virus L = 5: ities 7 5642	229E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (58%), Gaps = 13/478 (23  TYRRLISMMGFKMNYQVMGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY +IS MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+ TYEHVISYMGFRPDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV	66 5701
	Score Ident Query: Sbjct: Query:	virus = 5: ities 7 5642 67	229E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (28)  TYRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY +IS MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+ TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV  GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQM GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG FW+V+R +IVQM	66 5701 126
-	Score Ident: Query: Sbjct: Query:	virus L = 5: ities 7 5642 67 5702	229E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (29)  TYRRLISMMGFKMYYQVNGYPNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY + IS MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+ TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV  GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQM GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG PW+V+R +IVQM GFSNGVDFVAQPEGCVLTNTGSVVKFVRARAPPGEQFTHIVFLLRKGQPWSVLRKRIVQM	5701 126 5761
	Score Ident Query: Sbjct: Query: Sbjct: Query:	rirus L = 5: ities 7 5642 67 5702	229E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (28%)  TYRRLISMMGFKMNYQVNGYFNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL  TY + IS MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+  TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV  GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQM  GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG FW+V+R +IVQM  GFSNGVDFVAQPEGCVLTNTGSVVKPVRARAPFGEQFTHIVPLLRKGQFWSVLRKRIVQM  LSDTLKGLSDRVVFVLWAHGFELTSMKYFVKLGPERTCCLCDKRATCFSTSSDTYACWNH  ++D L G SD +VFVLWA G ELT+M+YFVKIG + C C ATC+++ S+ Y C++	5701 126 5761 186
-	Score Ident. Query: Sbjct: Query: Sbjct: Query: Sbjct:	rirus Long	229E]:  angth = 6758  35 bits (1379), Expect = e-151  = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (28)  TYRRLISMMGFKMNYQVNGYPNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL  TY + IS MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+  TYEHVISYMGFREDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV  GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLFWNVVRIKIVQM  GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG PW+V+R +IVQM  GFSNGVDFVAQPEGCVLTNTGSVVKPVRARAPPGEQFTHIVPLLRKGQPWSVLRKRIVQM  LSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGFERTCCLCDKRATCFSTSSDTYACWNH  ++D L G SD +VFVLWA G ELT+M+YFVKIG + C C ATC+++ S+ Y C+ H  IADFLAGSSDVLVFVLWAGGLELTTMRYFVKIGAVKH-CQCGTVATCYNSVSNDYCCFKH	5701 126 5761 186 5820
	Score Ident Query: Sbjct: Query: Sbjct: Query: Sbjct: Query:	rirus Long	225E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (28)  TYRRLISMMGFKMNYQVNGYPNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY + 1S MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+ TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV  GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPFGDQFKHLIPLMYKGLFWNVVRIKIVQM GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG PW+V+R +IVQM GFSMGVDFVAQPEGCVLTNTGSVVKFVRARAPFGEQFTHTVFLLRRGQFWSVLRKRIVQM  LSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGFFRTCCLCDKRATCFSTSSDTYACMNH ++D L G SD +VFVLWA G ELT+W+YFVKIG + C C ATC+++ S+ Y C+ H LADFLAGSSDVLVFVLWAHGGLELTTMRYFVKIGAVKH-CQCGTVATCYNSVSNDYCCFKH  SVGFDYVNNFFMIDVQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVKR ++G DYVYNP++ID+QQWGF G+L +NH C V H N HVAS DAIMTRCLAV++CFVKR	5) 66 5701 126 5761 186 5820 246
	Score Ident. Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Sbjct:	rirus Long	225E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (23)  TYRRLISMMGFKMNYQVNGYPNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY +IS MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQ+ TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV  GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQM GFS GV+ VA P G V T + V A+ PFG+QF H++PL+ KG PW+V+R +IVQM GFSNGVDFVAQPEGCVLTNTGSVVKPVRARAPPGEQFTHIVPLLRKGQFWSVLRKRIVQM LSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFFSTSDTYACWNH +HD L G SD +VPVLWA G ELT+M+YFVKIG + C C ATC+++ S+ Y C+ H LADFLAGSSDVLVFVLWAGGLELTTMRYFVKIGAVKH-CQCGTVATCYNSVSNDYCCFKH SVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVKR +HG DYVYNP++ID+QQWG+ G+L +NH C VH N HVAS DAIMTRCLAVHCFVK ALGCDYVYNPYVIDIQQWGFVGSLSTNHHAICNVHRNEHVASGDAIMTRCLAVHCFVK	5701 126 5761 186 5820 246 5880
	Score Ident. Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Query: Sbjct: Query:	247	229E]: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (28%); TYRRLISMMGFKMNYQVNGYPNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY +1S MGF+ + + G ++F TR+ A+RHVR W+G DUEG H T D VGTN+PLQ+ TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLPWNVVRIKIVQM GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG PW+V+R +IVQM GFSNGVDFVAQPEGCVLTNTGSVVKPVRARAPFGEQFTHIVPLLRKGQPWSVLRRIVQM LSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYACWNH ++D L G SD +VFVLWA G ELT+M+YFVKIG + C C ATC+++ S+ Y C+ H IADFLAGSSDVLVFVLWAGGLELTTMRYFVKIGAVKH-CQCGTVATCYNSVSNDYCCFKH SVGFDYVYNPFMIDVQQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHCFFVK ++G DYVYNP++ID+QQWG+ G+L +NH C VH N HVAS DAIMTRCLAV+CFVK ALGCDYVNPYVIDIQQWGYVGSLSTNHHAICNVHRNEHVASGDAIMTRCLAVYDCFVKN VDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEWK VDWSVEYPIIGDELRVNSACRKVQHMVVKSALLADKFPVLHDIGNPKAIKCVPQAEVEWK VDWSY YP+I +E +N R VQ +++++++	5701 126 5761 186 5820 246 5880 306
	Score Ident. Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Sbjct: Query: Sbjct: Sbjct:	ricus L.  = 55: ities  7 5642  67 5702  127 5762  187 5821  247 5881	225B: ength = 6758  35 bits (1379), Expect = e-151 = 254/478 (53%), Positives = 329/478 (68%), Gaps = 13/478 (28) TYRRLISMMGFKMNYQVNGYPNMFITREBAIRHVRAWIGFDVEGCHATRDAVGTNLPLQL TY + 15 MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQL TY + 15 MGF+ + + G ++F TR+ A+RHVR W+G DVEG H T D VGTN+PLQL TYEHVISYMGFRFDVSMPGSHSLFCTRDFAMRHVRGWLGMDVEGAHVTGDNVGTNVPLQV GFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDQFKHLIPLMYKGLFWNVVRIKIVQM GFS GV+ VA P G V T + V A+ PPG+QF H++PL+ KG PW+V+R +IVQM GFSNGVDFVAQPEGCVLTNTGSVVKFVRARAPPGEQFTHIVFLLRKGQFWSVLRKRIVQM LSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPETCCLCDKRATCFSTSSDTYACWNH ++D L G SD +VFVLWA G ELT+W+YFVKIG + C C ATC+++ S+ Y C+ H TADFLAGSSDVLVFVLWAHGGLELTTMRYFVKIGAVKH-CQCGTVATCYNSVSNDYCCFKH SVGFDYVYNPFMIDVQWGFTGNLQSNHDQHCQVHGNAHVASCDAIMTRCLAVHECFVKR +HG DYVYNP++ID+QQWG+ G+L +NH C VH N HVAS DAIMTRCLAV++CFVK ALGCDYVYNPYVIDIQQWGYVGSLSTNHHAICNVHRNEHVASGDAIMTRCLAVYDCFVKN VDWSVEYFIIGDELRVNSACRKVQHMVVKSALLADKFFVLHDIGNFKAIKCVPQAEVEWK	66 5701 126 5761 186 5820 246 5880 306 5939

```
YD P+
                         +E YY H
                                         DG+CLFWNCNVD YP +IVCRFDTR S L
 Sbjct: 5940 CYDKNPINSNVKTLE---YDYMTHGQ--MDGLCLFWNCNVDMYPEFSIVCRFDTRTRSTL 5994
 Ouery: 367 NLPGCDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSDSPCESHGKQVVSD-IDYVPL 425
             NL G +GGSLYVN HAFHTPA+DK A
                                       LK PFFYY D CE
                                                             VV D ++YVPI.
 Sbict: 5995 NLEGVNGGSLYVNNHAFHTPAYDKRAMAKLKPAPFFYYDDGSCE-----VVHDQVNYVPL 6049
Query: 426 KSATCITRCNLGGAVCRHHANEYRQYLDAYNMMISAGFSLWIYKQFDTYNLWNTFTRL 483
             ++ CIT+CN+GGAVC HAN YR Y+++YN+ AGF++W+
                                                         FD YNLW TFT +
 Sbict: 6050 RATNCITKCNIGGAVCSKHANLYRAYVESYNIFTOAGFNIWVPTTFDCYNLWOTFTEV 6107
 >gi|133591|sp|P18458|RRPB BEV
                                RNA-directed RNA polymerase (ORF1B)
 gi 94017 pir | S11238 polymerase - Berne virus
 gi | 1334814 | emb | CAA36601.1 |
                              2nd polymerase reading frame (AA 1-2291) [Berne
 virusl
          Length = 2291
 Score = 50.1 bits (118), Expect = 8e-05
 Identities = 37/103 (35%), Positives = 54/103 (52%), Gaps = 11/103 (10%)
Ouery: 140 FVLWAHGFELTSMKYFVKIGPERTC--CLCDKRATCFSTSSDTYACWNHSVGF--DYVYN 195
            F+L++
                  +L S+K++V+
                                   TC CC+AC
                                                     + Y C N
Sbict: 1511 FILYSCSNDLKSLKFYVEFD---TCYFCSCGEMAICLMRDGN-YKCRNCYGGMLISKLVN 1566
Query: 196 PFMIDVQQWGFTGNLQSNHDQHC-QVHGNAHVASCDAIMTRCL 237
               +DVO+
                         LO HD C Q HG++H A CDA+MT+CL
Sbjct: 1567 CKYLDVQKERV--KLQDAHDAICQQFHGDSHEALCDAVMTKCL 1607
>qi|1513061|dbj|BAA13323.1| cyanoprotein alpha subunit precursor [Riptortus
clavatus]
          Length = 693
 Score = 34.7 bits (78), Expect = 3.7
 Identities = 16/36 (44%), Positives = 22/36 (61%), Gaps = 1/36 (2%)
Query: 371 CDGGSLYVNKHAFHTPAFDKSAFTNLKQLPFFYYSD 406
           C G LY +KHA
                         P FD+ A+ + O+P FY+ D
Sbjct: 643 CGGSKLYDSKHAMGFP-FDRPAYPDAFQVPNFYFKD 677
  Database: All non-redundant GenBank CDS
  translations+PDB+SwissProt+PIR+PRF
    Posted date: Apr 11, 2003 2:30 AM
  Number of letters in database: 454,141,287
  Number of sequences in database: 1,411,415
```

Lambda K H 0.325 0.139 0.456

Gapped Lambda K H 0.267 0.0410 0.140

S2: 75 (33.5 bits)

#### 176/193

```
Matrix: BLOSUM62
Gap Penalties: Existence: 11, Extension: 1
Number of Hits to DB: 473.361.261
Number of Sequences: 1411415
Number of extensions: 20503315
Number of successful extensions: 51018
Number of sequences better than 10.0: 27
Number of HSP's better than 10.0 without gapping: 26
Number of HSP's successfully gapped in prelim test: 1
Number of HSP's that attempted gapping in prelim test: 50937
Number of HSP's gapped (non-prelim): 33
length of query: 486
length of database: 454,141,287
effective HSP length: 127
effective length of query: 359
effective length of database: 274,891,582
effective search space: 98686077938
effective search space used: 98686077938
T: 11
A: 40
x1: 15 (7.0 bits)
X2: 38 (14.6 bits)
X3: 64 (24.7 bits)
S1: 40 (21.6 bits)
```

FIGURE 125

# FIGURE 125A

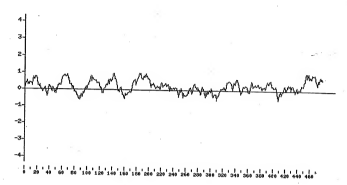
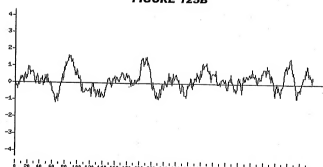


FIGURE 125B



#### FIGURE 126

#### 5'3' Frame 1

QVHQNVCVL-LIFYLMTLSR--SHKICQ-FQKWSRLQLTMLKFHSCFGVRMDMLKPSTQN YKQVKRGNQVLRCLTCTRCKECFLKSVTFRIMVKMLLYQKE---MSQSILNCVNT-IHLL -LYPPT-ELFTLVL

#### 5'3' Frame 2

RFIKMCVFCD-SFT--LCRDNKVTRFVSDFKSGQGYN-LC-NFIHALV-GWTC-NLLPKT TSKSSVATRCCDA-LVQDAKNAS-KV-PSELW-KCCYTKRNNDECRKVYSTVSILKYTYF SCTLOHESYSLWCW

#### 5'3' Frame 3

GSSKCVCSVIDLLLDDFVEIIKSQDLSVISKVVKVTIDYAEISFMLWCKDGHVETFYPKL QASQAWQPGVAMPNLYKMQRMLLEKCDLQNYGENAVIPKGIMMNVAKYTQLCQYLNTLTL AVPSNMRVIHFGAG

# 3'5' Frame 1

PAPK-ITLMLEGTAKVSVFKY-HS-VYFATFIIIPFGITAFSP-F-RSHFSRSILCILYK LGIATPGCHA-LACSFG-KVSTCPSLHQSMNEISA-SIVTLTTFEITDKSCDFIISTKSS SKRSITEHTHFDEP

#### 3'5' Frame 2

 $QHQSE-LSCWRVQLK-VYLSIDTVEYTLRHSSLFLLV-QHFHHNSEGHTFQEAFFASCTS\\ -ASQHLVATLDLLVVLGRRFQHVHPYTKA-MKFQHSQL-P-PLLKSLTNLVTLLSRQSHQVKDOSONTHILMNL\\$ 

#### 3'5' Frame 3

 ${\tt STKVNNSHVGGYS-SKCI-VLTQLSILCDIHHYSFWYNSIFTIILKVTLFKKHSLHLVQV RHRNTWLPRLTCL-FWVEGFNMSILTPKHE-NFSIVNCNLDHF-NH-QIL-LYYLDKVIK-KINHRTHTF-T$ 

#### FIGURE 127

## 5'3' Frame 1

-VFTYPGKANQPRSLVDLFSKRTN-NV--WTPIKPT-CPPHYIWWTHRFN-Q-PEWRTAM GQGQNSADPKVYPIILRLGSQLSLSMARRNLDSLEARAFQSTPIVVQMTKLATTEELPDE FVVVTAK-KSSAPDGTSIT-ELAQKLHFPTALTKKASYGLQLREP-IHPKTTLAPAILIT MLPPCYNFLKEQHCQKASTQREAEAAVKPLLAPHHVVAVIQEIQLLAAVGEILLLEWLAE VVKLPSRYCC-TD-TSLRAKFLVKANNNKAKLSLRNLLLRHLKSLAKNVLPQNSTTSLKH LGDVVQNKPKEISGTKT-SDKELITNIGPQIAQFA

# 5'3' Frame 2

RFLPTQEKPTNLDLL-ICSLNEQIKMSDNGPQSNQRSAPRITFGGPTDSTDNNQNGGLQW GKAKTAPTPRFTQ-YCVLVHSSHSAWQGGT-IPSRPGRSNQHQ-WSR-PNWLLPKSYPTS SWW-RQNERAQPQMVLLLPRNWPRSFTSLRR-QRRHRMGCN-GSLEYTQRPHWHPQS--Q CCHRATTSSRNNIAKRLLRRGKQRRQSSLFSLLIT-SR-FKKFNSWQQ-GKFSCSNG-RR W-NCPRAIAARQIEPA-EQSFW-RPTTTRPNCH-EICC-GI-KASPKTYCHKTVQRHSSI WETWSRTNPRKFRGPRPNQTRN-LQTLGRKLHNLP

## 5'3' Frame 3

GFYLPRKSQPTSISCRSVL-TNKLKCLIMDPNQTNVVPPALHLVDPQIQLTITRMEDCNG ARPKQRRPQGLPNNIASWFTALTQHGKEELRFPRGQGVPINTNSGPDDQIGYYRRATRRV RGGDGKMKELSPRWYFYYLGTGPEASLPYGANKEGIVWVATEGALNTPKDHIGTRNPNNN AATVLQLPQGTTLPKGFYAEGSRGGSQASSRSSRSRSRNSTPGSSRGNSPARMASGG GETALALLLLDRLNQLESKVSGKGQQQQGQTVTKKSAAEASKKPRQKRTATKQYNVTQAF GRRGPEQTQGNFGDQDLIRQGTDYKHWAANCTIC

# 3'5' Frame 1

RQIVQFAAQCL-SVPCLIRSWSPKFPWVCSGPRLPNA-VTLYCFVAVRFWRGFLDASAAD FLVTVWPCCCWPLPETLLSSWFNLSSSNSARAVSPPPLAIRAGEFPLLLPGVEFLELPRL RDEEREEA-LPPLLPSA-KPFGNVVP-GSCSTVAALLLGLRVPMWSLGVFKAPSVATHTM PSLLAP-GSEASGPVPR--KYHLGLSSFILPSPPRTRRVALR--PIWSSGPLLVLIGTPW PRGNLSSSLPC-VRAVNQDAILLGKPWGRRCFGLAPLQSSILVIVS-ICGSTKCNAGGTT LV-LGSIIRHFNLFV-RTDLQEIEVGWLFLGR-KP

#### 3'5' Frame 2

GKLCNLRPNVCNQFLV-LGLGPRNFLGFVLDHVSQMLE-RCTVLWQYVFGEAF-MPOOOI

S--QFGLVVVGLYQKLCSQAGSICLAAIARGQFHHLR-PFEQENFPYCCQELNFLNYRDY VMRSEKRLDCRLCFPLRRSLLAMLFLEEVVARWQHCY-DCGCQCGLWVYSRLPQLQPIRC LLC-RRREVKLLGQFLGNRSTIWG-ALSFCRHHHELVG-LFGSSQFGHLDHYWC-LERPG LEGI-VPPCHAE-EL-TKTQYYWVNLGVGAVLALPHCSPPFWLLSVESVGPPNVMRGALR WFDWGPLSDILICSFREQIYKRSRLVGFSWVGKNL

# 3'5' Frame 3

ANCAICGPMFVISSLSD-VLVPEISLGLFWTTSPKCLSDVVLFCGSTFLARLFRCLSSRF LSDSLALLLLAFTRNFALKLVQSV-QQ-REGSFTTSASHSSRRISPTAARS-IS-ITATT --GARRGLTAASASLCVEAFWQCCSLRKL-HGGSIVIRIAGANVVFGCIQGSLSCNPYDA FFVSAVGK-SFWASS-VIEVPSGAELFHFAVTTTNSSGSSSVVANLVIWTTIGVDWNALA SRESKFLLAMLSESCEPRRNIIG-TLGSALFWPCPIAVLHSGYCQLNLWVHQM-CGGHYV GLIGVHYQTF-FVRLENRSTRDRGWLAFPG-VKT

## FIGURE 128

-GLELKL-LTSICAF-PFCYSLF--CLLYFGFHSKSRI-KNLVPKSKRT-NFSLF-LVFL
YAVAYAL-YSAVHLINLMCLKILVRYNTRGNTYSTAWLCALGKVLPFHRWHTMVQTCTPN
VTINCQDPAGGALIARCWYLHEGHQTAAFRDVLVVLNKRTN-NV--WTPIKPT-CPPHYI
WWTHRFN-Q-PEWRTQWGKAKTAPTPRFTQ-YCVLVHSSHSAWQGGT-IPSRPGRSNQHQ
-WSR-PNWLLPKSYPTSSW-RQNERAQPQMVLLLPRNWPRSFTSLRF-QRRHRMGCN-G
SLEYTQRPHWHPQS--QCCHRATTSSRNNIAKRLLRRGKQRRQSSLFSLLTI-SR-FKKF
NSWQQ-GKFSCSNG-RRW-NCPRAIAARQIEPA-EQSFW-RPTTTRPNCH-EICC-GI-K
ASPKYTCHKTVQRHSSIWETWSRTNPRKFRGPRPNQTRN-LQTLAANCTICSKCLCILWN
VTHWHGSHTFGNMADLSWSH-IG-QRSTIQRQRHTAEQAH-RIQNIPTNRA-KGQKEKD--SSAFAAETKEAAHCDSSSC

EDSSSSFN-LLFVLFSLSAIPCFNNAYYILVFTRNPGSRRTLYQSLNEHETSHCFDLYFS MQLHMHCSTALCI--TSCA-RSL-GTTLGVILIALLGFVL-ERFYLFIDGTLWFKHAHLM LLSTVKIQLVVRL-LGVGTFMKVTKLLHLETYLLF-INEQIKMSDNGPQSNQRSAPRITF GGPTDSTDNNQNGGRNGARPKQRRPQGLPNNIASWFTALTQHGKEELRFPRGQGVPINTN SGPDDQIGYYRRATRRVRGGDGKMKELSPRWFFYYLGTGPEASLFYGANKEGIVWVATEG ALNTPKDHIGTRNPNNNAATVLQLPQGTTLPKGFYAEGSRGSQSSSSSSSSSRSNSNS TPGSSRGNSPARMASGGETALALLLDRLNQLESKVSGKGQQQQGQTVTKKSAAEASKK PRQKRTATKQYNVTQAFGRRGPEQTQGNFGDQDLIRQGTDYKHWPQIAQFAPSASAFFGM SRIGMEVTPSGTWLTYHGAIKLDDKDPQFKDNVILLNKHIDAYKTFPPTEPKKDKKKKTD EAOPLPOROKKOPTVTLLP

RTRAQALIDFYLCFLAFLLFLVLIMLIIFWFSLEIQDLEEPCTKV-TNMKLLIVLTCISL CSCICTVVQRCASNKPHVLEDPCKVQH-G-YL-HCLALCSRKGFTFS-MAHYGSNMHT-C YYQLSRSSWWCAYS-VLVPS-RSPNCCI-RRTCCFK-TNKLKCLIMDPNQTNVVPPALHL VDPQIQLITITMEDAMGQGQNSADPKVYPIILRLGSQLSLSMARRNLDSLEARAFQSTPI VVQMTKLATTEELPDEFVVVTAK-KSSAPDGTSIT-ELAQKLHFPTALTKKASYGLQLRE P-IHPKTTLAPAILITMLPPCYNFLKEQHCQKASTQRBAEAAVKPLLAPHHVVAVIQEIQ LLAAVGEILLLEWLAEVVKLPSRYCC-TD-TSLRAKFLVKANNNKAKLSLRNLLLRHLKS LAKNVLPQNSTTSLKHLGDVVQNKPKEISGTKT-SDKELITNIGRKLHNLLQVPLHSLEC HALAWKSHLREHG-LIMEPLNWMTKIHNSKTTSYC-TSTLTHTKHSHQQSLKRTKRKRLM KLSLCRRDKRSSPL-LFFL

#### FIGURE 129

# 5'3' Frame 1

#### 5'3' Frame 2

# 5'3' Frame 3

## 3'5' Frame 1

#### 3'5' Frame 2

# 3'5' Frame 3

# FIGURE 130 20 40 50 60 SEO ID NO: 9997 KGHDLRRLISMMGFKMNYQVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVG SEO ID NO:10034 ----YRRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLP Prim. Cons. KGHD2RRLISMMGFKMNYOVNGYPNMFITREEAIRHVRAWIGFDVEGCHATRDAVGTNLP 70 80 100 SEQ ID NO: 9997 LQLGFSTGVNLVAVPTGYVDTENNTKFTRVNAQTSTSEQFKHLIPLMYKGLPWNVVRIKI SEQ ID NO:10034 LQLGFSTGVNLVAVPTGYVDTENNTEFTRVNAKPPPGDOFKHLI----********************* Prim. cons. LQLGFSTGVNLVAVPTGYVDTENNT2FTRVNA222222OFKHLIPLMYKGLPWNVVRIKI 130 140 150 160 170 180 SEQ ID NO: 9997 VQMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFS Prim. cons. ${\tt VQMLSDTLKGLSDRVVFVLWAHGFELTSMKYFVKIGPERTCCLCDKRATCFSTSSDTYAC}$ 190 200

SEQ ID NO: 9997 WNHSVGFDYVYNPFMIDVQQWGLYG

# FIGURE 131

# 5'3' Frame 1

# 5'3' Frame 2

 $\begin{array}{c} {\rm caggttcatcaaaaatgtgtgttctgtgattgattgatctttactgtgatgacttttgtcgagat} \\ R \ F \ I \ K \ M \ C \ V \ F \ C \ D \ - \ S \ F \ T \ - \ L \ C \ R \ D \\ {\rm aataaagtcacaagatttgtcagtgatttcaaaagtggtcaaagttacaattgactattg} \\ N \ K \ V \ T \ R \ F \ V \ S \ D \ F \ K \ S \ G \ Q \ G \ Y \ N \ - \ L \ C \\ {\rm tgaaatttcattcattcatctttggtgtaaaggattgtcgaaccttctacccaaaact} \\ - \ N \ F \ I \ H \ A \ L \ V \ - G \ W \ T \ C \ - \ N \ L \ L \ P \ K \ T \\ {\rm acaagcaagtcaagtgtgcaaccaggtgttcgaatgcctaacttgtacaaggtgcaaag} \\ T \ S \ K \ S \ S \ V \ A \ T \ R \ C \ C \ D \ A \ - \ L \ V \ Q \ D \ A \ K \\ {\rm aatgcttcttgaaaagtgtgaccttcaacttgtacaaagg} \\ T \ S \ K \ S \ S \ V \ A \ T \ R \ C \ C \ D \ A \ - \ L \ V \ Q \ D \ A \ K \\ {\rm aatgcttcttgaaaagtgtgaccttcaacttgtgaaaatgctgttataccaaaagg} \\ N \ A \ S \ - \ K \ V \ - \ P \ S \ E \ L \ W \ - \ K \ C \ C \ Y \ T \ K \ R \\ {\rm aataatatgatgaatgtcgcaaagtatacctcaacttgtgtcaatactttaaataccttacttt} \\ N \ N \ D \ E \ C \ R \ K \ V \ Y \ S \ T \ V \ S \ I \ L \ K \ Y \ T \ Y \ F \\ {\rm agctgtaccctccaacatgaggattattcactttggtcgc} \\ S \ C \ T \ L \ O \ H \ E \ S \ Y \ S \ L \ W \ C \ W \end{array}$ 

# 5'3' Frame 3

caggttcatcaaaatgtgtgtgttctgtgattgatcttttacttgatgactttgtcgagata G S S K C V C S V I D L L L D D F V E I ataaagtcacaagatttgtcagtgatttcaaaagtggtcaaggttacaattgactatgct I K S Q D L S V I S K V V K V T I D Y A gaaatttcattcatgcttggtgtaaggatggacatgttgaaaccttctacccaaaacta E I S F M L W C K D G H V E T F Y P K L caagcaagtcaagcttgtgcaaccagtgttgcgatgcctaacttgtacaagatgcaaaga

Q A S Q A W Q P G V A M P N L Y K M Q R atgettettgaaaagtgtgacetteagaattatggtgaaaatgetgttataceaaaagga M L L E K C D L Q N Y G E N A V I P K G ataatgatgaatgtegeaagtatacteaactgtgtcaatacttaaatacacttacttta I M M N V A K Y T Q L C Q Y L N T L T L getgtacectceaacatgagagttattcactttggtgetgg A V P S N M R V I H F G A

# 3'5' Frame 1

# 3'5' Frame 2

3'5' Frame 3

# FIGURE 132

### 5'3' Frame 1

2004092360A2 1 3

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# 5'3' Frame 2

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#### 5'3' Frame 3

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#### 3'5' Frame 1

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LYCFVAVRFWRGFLDASAAD ttettagtgacagtttggeettgttgttgttggeetttaecagaaaetttgetetcaage F L V T V W P C C C W P L P E T L L S S tggttcaatctgtctagcagcaatagcqcgagggcagtttcaccacctccgctagccatt W F N L S S S N S A R A V S P P P L A I cgagcaggagaatttcccctactgctgccaggagttgaatttcttgaattaccgcgacta RAGEFPLLLPGVEFLELPRL cgtgatgaggagcgagaagaggcttgactgccqcctctgcttccctctgcqtagaagcct R D E E R E E A - L P P L L P S A - K P tttggcaatgttgttccttgaggaagttgtagcacggtggcagcattgttattaggattg F G N V V P - G S C S T V A A L L L G L cqggtqccaatgtqqtctttqqqtgtattcaaggctccctcagttgcaacccatacqatq RVPMWSLGVFKAPSVATHTM ccttctttgttagcgccgtagggaagtgaagcttctgggccagttcctaggtaatagaag PSLLAP-GSEASGPVPR--K taccatctggggctgagctctttcattttgccgtcaccaccacgaactcgtcgggtagct Y H L G L S S F I L P S P P R T R R V A cttcggtagtagccaatttggtcatctggaccactattggtgttgattggaacqccctgg LR--PIWSSGPLLVLIGTPW PRGNLSSSLPC-VRAVNODA atattattgggtaaaccttggggtcggcgctgttttggccttgccccattgcagtcctcc ... I L L G K P W G R R C F G L A P L O S S  ${\tt attctqgttattqtcaqttqaatctgtqggtccaccaaatgtaatgcggggggcactacg}$ ILVIVS-ICGSTKCNAGGTT ttggtttgattggggtccattatcagacattttaatttgttcgtttagagaacagatcta LV-LGSIIRHFNLFV-RTDL caagagatcgaggttggttggcttttcctgggtaggtaaaaaccta Q E I E V G W L F L G R - K P

# 3'5' Frame 2

CID: <WO____2004092360A2_[_>

aggcaaattgtgcaatttgcggcccaatgtttgtaatcagttccttgtctgattaggtctt G K L C N L R P N V C N Q F L V - L G L ggtccccgaaatttccttgggtttgttctggaccacgtctcccaaatgcttgagtgacgt G P R N F L G F V L D H V S Q M L E - R tgtactgttttgtggcagtacgtttttgggcgtttttagatgcctcagcagagtt C T V L W Q Y V F G E A F - M P Q Q Q I tcttagtgacagtttggccttgttgttgttggcctttaccagaaactttgctctcaagct S - - Q F G L V V V G L Y Q K L C S Q A ggttcaatctgtctgcagcagtagcgagtggaagttcaccacctccgctagccattc G S I C L A A I A R G Q F H H L R - P F gagcagagaatttcccctactgctgccaggagttgaatttcttgaattaccggactacc E Q E N F P Y C C Q E L N F L N Y R D Y gtgatgaggagagagagagagagagttgactgccctctcgcttcgctgcagagacct V M R S E K R L D C R L C F P L R R S L ttggcaatgttgtccttgagagaagttgt

TAMLFLEEVVARWOHCY-DC gggtgccaatgtggtctttgggtgtattcaaggctccctcagttgcaacccatacgatgc GCOCGLWVYSRLPQLOPIRC cttctttgttagcgccgtagggaagtgaagcttctgggccagttcctaggtaatagaagt LLC-RRREVKLLGOFLGNRS accatctggggctgagctctttcattttgccgtcaccaccacgaactcgtcgggtagctc TIWG-ALSFCRHHHELVG-L ttcggtagtagccaatttggtcatctggaccactattggtgttgattggaacgccctggc FGSSOFGHLDHYWC-LERPG I. E.G. I - V P P C H A E - E L - T K T O tattattgggtaaaccttggggtcggcqctgttttggccttqccccattqcagtcctcca YYWVNLGVGAVLALPHCSPP ttctggttattgtcagttgaatctgtgggtccaccaaatgtaatgcggggggcactacgt FWLLSVESVGPPNVMRGALR taatttaattaagatccattatcagacattttaatttattattaagagaacagatctac W F D W G P L S D I L I C S F R E O I Y aagagatcgaggttggttggcttttcctgggtaggtaaaaaccta K. R. S. R. L. V. G. F. S. W. V. G. K. N. L.

## 3'5' Frame 3

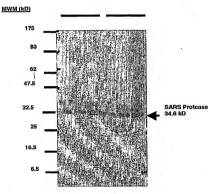
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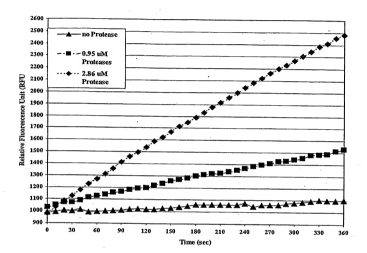
S R E S K F L L A M L S E S C E P R R N attattgggtaaaccttggggcgctgttttggccttgcccattgcagtcccat I I G - T L G S A L F W P C P I A V L H tctggttattgtcagttgaatctgtgggtccaccaaatgtaatgcgggggggcactgtt S G Y C Q L N L W V H Q M - C G G H Y V ggtttgattggggtccattatcagacatttaattgttcgtttagagaacagatctaca G L I G V H Y Q T F - F V R L E N R S T agagatcgaggtggtggtggttgctttcctgggtaggtaaaaccca R D R G W L A F P G - V K T

FIGURE 133





# FIGURE 134



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Jens-Peter [DE/DE]; c/o Chiron Corporation, P.O.

Box 8097, Emeryville, CA 94662-8097 (US), CHIEN. David [US/US]; c/o Chiron Corporation, P.O. Box 8097,

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Emeryville, CA 94662-8097 (US). HAN, Jang [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). POLO, John [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). WEINER, Amy [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). HOUGHTON, Michael [GB/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). SONG, Hyun, Chul [KR/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). SEO. Mi, Young [KR/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). DONNELLY, John, J. [US/US]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). KLENK, Hans, Dieter [DE/DE]; c/o Chiron Corporation, P.O. Box

8097, Emeryville, CA 94662-8097 (US). VALIANTE,

Nicholas [US/US]; c/o Chiron Corporation, P.O. Box

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- (71) Applicant (for all designated States except US): CHI-RON CORPORATION [US/US]; 4560 Horton Street, Emeryville, CA 94608-2916 (US).
- (72) Inventors; and

(75) Inventors/Applicants (for US only): RAPPUOLL. Rino [IT/IT]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). MASIGNANI, Vega [IT/IT]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). STADLER, Konrad [DE/DE]; c/o Chiron Corporation, P.O. Box 8097, Emeryville, CA 94662-8097 (US). GREGERSEN,

(74) Agents: HALE, Rebecca, M. et al.; Chiron Corporation, Intellectual Property R338, P.O. Box 8097, Emeryville, CA 94662-8097 (US).

8097, Emeryville, CA 94662-8097 (US).

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[Continued on next page]

(54) Title: THE SEVERE ACUTE RESPIRATORY SYNDROME CORONAVIRUS

(57) Abstract: An outbreak of a virulent respiratory virus, now known as Severe Acute Respiratory Syndrome (SARS), was identified in Hong Kong, China and a growing number of countries around the world in 2003. The invention relates to nucleic acids and proteins from the SARS coronavirus. These nucleic acids and proteins can be used in the preparation and manufacture of vaccine formulations, diagnostic reagents, kits, etc. The invention also provides methods for treating SARS by administering small molecule antiviral compounds, as well as methods of identifying potent small molecules for the treatment of SARS.

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- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments
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A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C07K14/165 C120 C1201/70 C07K16/10 A61K35/76 A61K38/16 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system tollowed by classification symbols) IPC 7 CO7K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the International search (name of data base and, where practical, search terms used) EPO-Internal, BIOSIS, Sequence Search, EMBASE, PAJ, WPI Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Category * Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. PEIRIS J S M ET AL: "Coronavirus as a 1-28. possible cause of severe acute respiratory 73-77, syndrome." 79,83, LANCET. 19 APR 2003. 85-109 vol. 361, no. 9366, 8 April 2003 (2003-04-08), pages 114-120 1319-1325, XP004421148 ISSN: 0140-6736 cited in the application published online on 8 April 2003 (http://image.thelancet.com/extras/03art34 77web.pdf) page 1320, left-hand column - page 1321, right-hand column page 1322, right-hand column page 1322, left-hand column; figure 3 -/--X Further documents are listed in the continuation of box C. Y Patent family members are listed in annex. Special categories of cited documents: *T* later document published after the International tiling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the an which is not considered to be of particular relevance earlier document but published on or after the international *X* document of particular relevance; the claimed Invention cannot be considered novel or cannot be considered to Involve an inventive step when the document is taken alor filing date *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled in the art. O document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed *&* document member of the same patent tamily Date of the actual completion of the international search Date of mailing of the international search report 1 March 2005 n 8. 06. 2005 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016

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D	
DOX II	Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)
This trite	ornational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
	Although claims $77-83$ , $117-120$ are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
з. 🗍	Claims Nos.:
	because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box III	Observations where unity of invention is lacking (Continuation of item 3 of first sheet)
This Inte	rnational Searching Authority found multiple Inventions in this international application, as follows:
3	
	see additional sheet
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims,
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. [	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
. 🗀	
4. LX.J	No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the Invention first mentioned in the claims; it is covered by claims Nos.:
-	1-12(partially), 13(partially), 14, 15, 16-21(partially), 22-28(partially) 73-77 (partially), 79(partially), 83(partially), 85-93(partially), 94-98 99-104(partially), 105, 106-107(partially), 108, 109, 114-120(partially)
Remark	on Protest The additional search fees were accompanied by the applicant's protest.
	No protest accompanied the payment of additional search fees.

# FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

#### Invention 1

claims: 1-12(partially), 13(partially), 14, 15, 16-21(partially), 22-28(partially), 73-77(partially), 79(partially), 85-93(partially), 98-94(partially), 98-94(partially), 98-94(partially), 98, 109, 114-120(partially), 105, 106-107(partially), 108, 109, 114-120(partially)
The polypeptide is a spike (S) polypeptide encoded by SEQ. ID n 6042, nucleic acid encoding the spike protein, fragments thereof, antibodies specific for the spike protein, immunoassays using these antibodies, a vaccine comprising a spike protein, a viral vector comprising the protein, an immunogenic fragment thereof, double stranded RNA thereof, the recombinaint expression thereof and the medical use thereof.

#### Invention 2

claims:1-12(partially),13(partially),16-21(partially), 22-28(partially), 73-77(partially), 79(partially), 83(partially), 85-93(partially), 99-104(partially), 106-107(partially), 114-120(partially) the polypeptide is an envelope (E) polypeptide encoded by SEO. ID n 6045, nucleic acid encoding the E protein, fragments thereof, antibodies specific for the E protein, immunoassays using that antibody, a vaccine comprising an E protein, a viral vector comprising the protein,double stranded RNA thereof the recombinant expression thereof and the medical use thereof.

#### Invention 3

claims: 1-12(partially), 13(partially), 16-21(partially), 22-28(partially), 73-77(partially), 79(partially), 83(partially), 89-3(partially), 99-104(partially), 106-107(partially), 114-120(partially), 105-107(partially), 114-120(partially), 106-107(partially), 114-120(partially), 106-107(partially), 106-107(partially), 106-107(partially), 106-107(partially), 106-107(partially), 106-107(partially), 107(partially)

#### Invention 4

# FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Claims :1-12(partially), 13(partially), 16-21(partially), 22-27(partially), 73-7(partially), 79(partially), 85-93(partially), 99-104(partially), 106-107(partially), 114-120(partially), 106-107(partially), 114-120(partially) the polypeptide is a hemagglutinin esterase (HE) polypeptide, nucleic acid encoding the HE protein, fragment thereof, antibodies specific for the HE protein, immunoassays using that antibody, a vaccine comprising a HE protein, a viral vector comprising the protein, double stranded RNA thereof, the recombinant expression thereof and the medical use thereof.

# Invention 5

claims:1-12(partially),13(partially),16-21(partially), 22-28(partially), 73-77(partially), 79(partially), 83(partially), 89-93(partially), 99-104(partially), 106-107(partially), 114-120(partially), 106-107(partially), 114-120(partially) polypeptide encoded by SEQ. ID n 6052, nucleic acid encoding the N protein, fragment thereof, antibodies specific for the N protein, immunoassays using that antibody, a vaccine comprising a N protein, a viral vector comprising the protein, double stranded RNA thereof, the recombinant expression thereof and the medical use thereof.

# Invention 6

claims:1-12(partially);13(partially),16-21(partially), 22-27(partially), 73-77(partially), 79(partially), 83(partially), 89-93(partially), 99-104(partially), 106-107(partially), 114-120(partially)
The ORFIa polypeptide encoded by SEQ. ID n 6039, the preoteolytic fragments thereof such as NSP1-Nsp-7 corresponding to SEQ.ID n 9766-9774), nucleic acid encoding these proteins, fragments thereof, antibodies specific for these proteins, immunoassays using these antibodies, a vaccine comprising the protein, a viral vector comprising the protein, double stranded RNA thereof, the recombinant expression thereof and the medical use thereof.

Invention 7

# FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

# Invention 8

claims: 22(partially), 29-58, 84, 110-113(partially): 110-113(partially), 114-118(partially): A vaccine comprising an inactivated/attenuated SARS virus, a method of inactivating the SARS virus, a method of making an inactivated SARS vaccine

# Invention 9:

claims: 77(partially), 78, 79(partially), 80-82, 83 (patially), 119-120 (partially) A method of treatment of a patient suffering from SARS and a method of identifying a therapeutically active agent comprising measuring the attenuation of a SARS related enzyme, a method of treatment using a therapeutical agent of claims 77-82.

# Invention 10-7760

claims :59-72(partially)
A single-stranded oligonucleotide selected from the group consisting of the SEQ. IDs 21-6020, 6076-6568, 6586-6587, 7292-7301, 7325-7328, 7332-7352, 7353-7385, 10235-10298, 10352-10504, 10580-11322, 11325-11551 (taken from the list of claim number 59), PCR ktt comprising these primers, a method of detecting the presence of SARS virus in a sample using PCR.